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SINGLE NUCLEOTIDE POLYMORPHISMS IN GENES

RELATED APPLICATIONS

This application claims the benefit of U.S. Provisional Application Serial No. 60/153,357, filed September 10, 2000, U.S. Provisional Application Serial No. 60/220,947, filed July 26, 2000, and U.S. Provisional Application Serial No. 60/225,724, filed August 16, 2000, the entire teachings of all of which are incorporated herein by reference.

BACKGROUND OF THE INVENTION

The genomes of all organisms undergo spontaneous mutation in the course of
their continuing evolution, generating variant forms of progenitor nucleic acid
sequences (Gusella, Ann. Rev. Biochem. 55, 831-854 (1986)). The variant form may
confer an evolutionary advantage or disadvantage relative to a progenitor form, or may
be neutral. In some instances, a variant form confers a lethal disadvantage and is not
transmitted to subsequent generations of the organism. In other instances, a variant
form confers an evolutionary advantage to the species and is eventually incorporated
into the DNA of many or most members of the species and effectively becomes the
progenitor form. In many instances, both progenitor and variant form(s) survive and co-

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exist in a species population. The coexistence of multiple forms of a sequence gives rise to polymorphisms.

Several different types of polymorphism have been reported. A restriction fragment length polymorphism (RFLP) is a variation in DNA sequence that alters the length of a restriction fragment (Botstein *et al.*, *Am. J. Hum. Genet.* 32, 314-331 (1980)). The restriction fragment length polymorphism may create or delete a restriction site, thus changing the length of the restriction fragment. RFLPs have been widely used in human and animal genetic analyses (see WO 90/13668; W090/11369; Donis-Keller, *Cell 51*, 319-337 (1987); Lander *et al.*, *Genetics 121*, 85-99 (1989)).

When a heritable trait can be linked to a particular RFLP, the presence of the RFLP in an individual can be used to predict the likelihood that the animal will also exhibit the trait.

Other polymorphisms take the form of short tandem repeats (STRs) that include tandem di-, tri- and tetra-nucleotide repeated motifs. These tandem repeats are also referred to as variable number tandem repeat (VNTR) polymorphisms. VNTRs have been used in identity and paternity analysis (US 5,075,217; Armour *et al.*, *FEBS Lett.* 307, 113-115 (1992); Horn *et al.*, W0 91/14003; Jeffreys, EP 370,719), and in a large number of genetic mapping studies.

Other polymorphisms take the form of single nucleotide variations between individuals of the same species. Such polymorphisms are far more frequent than RFLPs, STRs and VNTRs. Some single nucleotide polymorphisms (SNP) occur in protein-coding nucleic acid sequences (coding sequence SNP (cSNP)), in which case, one of the polymorphic forms may give rise to the expression of a defective or otherwise variant protein and, potentially, a genetic disease. Examples of genes in which polymorphisms within coding sequences give rise to genetic disease include β -globin (sickle cell anemia), apoE4 (Alzheimer's Disease), Factor V Leiden (thrombosis), and CFTR (cystic fibrosis). cSNPs can alter the codon sequence of the gene and therefore specify an alternative amino acid. Such changes are called "missense" when another amino acid is substituted, and "nonsense" when the alternative codon specifies a

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stop signal in protein translation. When the cSNP does not alter the amino acid specified the cSNP is called "silent".

Other single nucleotide polymorphisms occur in noncoding regions. Some of these polymorphisms may also result in defective protein expression (e.g., as a result of defective splicing). Other single nucleotide polymorphisms have no phenotypic effects.

Single nucleotide polymorphisms can be used in the same manner as RFLPs and VNTRs, but offer several advantages. Single nucleotide polymorphisms occur with greater frequency and are spaced more uniformly throughout the genome than other forms of polymorphism. The greater frequency and uniformity of single nucleotide polymorphisms means that there is a greater probability that such a polymorphism will be found in close proximity to a genetic locus of interest than would be the case for other polymorphisms. The different forms of characterized single nucleotide polymorphisms are often easier to distinguish than other types of polymorphism (e.g., by use of assays employing allele-specific hybridization probes or primers).

Only a small percentage of the total repository of polymorphisms in humans and other organisms has been identified. The limited number of polymorphisms identified to date is due to the large amount of work required for their detection by conventional methods. For example, a conventional approach to identifying polymorphisms might be to sequence the same stretch of DNA in a population of individuals by dideoxy sequencing. In this type of approach, the amount of work increases in proportion to both the length of sequence and the number of individuals in a population and becomes impractical for large stretches of DNA or large numbers of persons.

SUMMARY OF THE INVENTION

Work described herein pertains to the identification of polymorphisms which can predispose individuals to disease, by resequencing large numbers of genes in a large number of individuals. Various genes from a number of individuals have been resequenced as described herein, and SNPs in these genes have been discovered (see the Table and Fig. 3). Some of these SNPs are cSNPs which specify a different amino acid

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sequence, some of the SNPs are silent cSNPs and some of these cSNPs specify a stop signal in protein translation. Some of the identified SNPs were located in non-coding regions.

The invention relates to a gene which comprises a single nucleotide polymorphism at a specific location. In a particular embodiment the invention relates to the variant allele of a gene having a single nucleotide polymorphism, which variant allele differs from a reference allele by one nucleotide at the site(s) identified in the Table and Fig. 3. Complements of these nucleic acid sequences are also included. The nucleic acid molecules can be DNA or RNA, and can be double- or single-stranded.

10 Nucleic acid molecules can be, for example, 5-10, 5-15, 10-20, 5-25, 10-30, 10-50 or 10-100 bases long.

The invention further provides allele-specific oligonucleotides that hybridize to the reference or variant allele of a gene comprising a single nucleotide polymorphism or to the complement thereof. These oligonucleotides can be probes or primers.

The invention further provides a method of analyzing a nucleic acid from an individual. The method determines which base is present at any one of the polymorphic sites shown in the Table and/or Fig. 3. Optionally, a set of bases occupying a set of the polymorphic sites shown in the Table and /or Fig. 3 is determined. This type of analysis can be performed on a number of individuals, who are tested for the presence of a disease phenotype. The presence or absence of disease phenotype is then correlated with a base or set of bases present at the polymorphic site or sites in the individuals tested.

Thus, the invention further relates to a method of predicting the presence, absence, likelihood of the presence or absence, or severity of a particular phenotype or disorder associated with a particular genotype. The method comprises obtaining a nucleic acid sample from an individual and determining the identity of one or more bases (nucleotides) at polymorphic sites of genes described herein, wherein the presence of a particular base is correlated with a specified phenotype or disorder, thereby

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predicting the presence, absence, likelihood of the presence or absence, or severity of the phenotype or disorder in the individual.

The thrombospondins are a family of extracellular matrix (ECM) glycoproteins

that modulate many cell behaviors including adhesion, migration, and proliferation. Thrombospondins (also known as thrombin sensitive proteins or TSPs) are large molecular weight glycoproteins composed of three identical disulfide-linked polypeptide chains. The results described herein also reveal an important association between alterations, particularly SNPs, in TSP genes, particularly TSP-1 and TSP-4, and vascular disease. In particular, SNPs in these genes which are associated with premature coronary artery disease (CAD)(or coronary heart disease) and myocardial infarction (MI) have been identified and represent a potentially vital marker of upstream biology influencing the complex process of atherosclerotic plaque generation and vulnerability.

Thus, the invention relates to the TSP gene SNPs identified as described herein, both singly and in combination, as well as to the use of these SNPs, and others in TSP genes, particularly those nearby in linkage disequilibrium with these SNPs, for diagnosis, prediction of clinical course and treatment response for vascular disease, development of new treatments for vascular disease based upon comparison of the variant and normal versions of the gene or gene product, and development of cell-culture based and animal models for research and treatment of vascular disease. The invention further relates to novel compounds and pharmaceutical compositions for use in the diagnosis and treatment of such disorders. In preferred embodiments, the vascular disease is CAD or MI.

The invention relates to isolated nucleic acid molecules comprising all or a portion of the variant allele of TSP-1 (e.g., as exemplified by SEQ ID NO: 1), and to isolated nucleic acid molecules comprising all or a portion of the variant allele of TSP-4 (e.g., as exemplified by SEQ ID NO: 3). Preferred portions are at least 10 contiguous nucleotides and comprise the polymorphic site, e.g., a portion of SEQ ID NO: 1 which is at least 10 contiguous nucleotides and comprises the "G" at position 2210, or a

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portion of SEQ ID NO: 3 which is at least 10 contiguous nucleotides and comprises the "C" at position 1186. The invention further relates to isolated gene products, e.g., polypeptides or proteins, which are encoded by a nucleic acid molecule comprising all or a portion of the variant allele of TSP-1 or TSP-4 (e.g., SEQ ID NO: 1 or SEQ ID NO: 3, respectively). The invention also relates to nucleic acid molecules which hybridize to and/or share identity with the variant alleles identified herein (or their complements) and which also comprise the variant nucleotide at the SNP site.

The invention further relates to isolated proteins or polypeptides comprising all or a portion of the variant amino acid sequence of TSP-1 (e.g., as exemplified by SEQ ID NO: 2), and to isolated proteins or polypeptides comprising all or a portion of the variant amino acid sequence of TSP-4 (e.g., as exemplified by SEQ ID NO: 4). Preferred polypeptides are at least 10 contiguous amino acids and comprise the polymorphic amino acid, e.g., a portion of SEQ ID NO: 2 which is at least 10 contiguous amino acids and comprises the serine at residue 700, or a portion of SEQ ID NO: 4 which is at least 10 contiguous amino acids and comprises the proline at residue 387. The invention further relates to isolated nucleic acid molecules encoding such proteins and polypeptides, as well as to antibodies which bind, e.g., specifically, to such proteins and polypeptides.

The invention further relates to a method of diagnosing or aiding in the diagnosis of a disorder associated with the presence of one or more of (a) a G at nucleotide position 2210 of SEQ ID NO: 1; or (b) a C at nucleotide position 1186 of SEQ ID NO: 3 in an individual. The method comprises obtaining a nucleic acid sample from the individual and determining the nucleotide present at one or more of the indicated nucleotide positions, wherein presence of one or more of (a) a G at nucleotide position 2210 of SEQ ID NO: 1; or (b) a C at nucleotide position 1186 of SEQ ID NO: 3 is indicative of increased likelihood of said disorder in the individual as compared with an appropriate control, *e.g.*, an individual having the reference nucleotide at one or more of said positions. In a particular embodiment the disorder is a vascular disease selected from the group consisting of atherosclerosis, coronary heart or artery disease, MI,

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stroke, peripheral vascular diseases, venous thromboembolism and pulmonary embolism. In a preferred embodiment, the vascular disease is selected from the group consisting of CAD and MI.

The invention further relates to a method of diagnosing or aiding in the diagnosis of a disorder associated with one or more of (a) a G at nucleotide position 2210 of SEQ ID NO: 1; or (b) a C at nucleotide position 1186 of SEQ ID NO: 3 in an individual. The method comprises obtaining a nucleic acid sample from the individual and determining the nucleotide present at one or more of the indicated nucleotide positions, wherein presence of one or more of (a) an A at nucleotide position 2210 of SEQ ID NO: 1; or (b) a G at nucleotide position 1186 of SEQ ID NO: 3 is indicative of decreased likelihood of said disorder in the individual as compared with an appropriate control, *e.g.*, an individual having the variant nucleotide at said position. In a particular embodiment the disorder is a vascular disease selected from the group consisting of atherosclerosis, coronary heart or artery disease, MI, stroke, peripheral vascular diseases, venous thromboembolism and pulmonary embolism. In a preferred embodiment, the vascular disease is selected from the group consisting of CAD and MI.

In one embodiment, the invention relates to a method for predicting the likelihood that an individual will have a vascular disease (or aiding in the diagnosis of a vascular disease), comprising the steps of obtaining a DNA sample from an individual to be assessed and determining the nucleotide present at one or more of nucleotide positions 2210 of SEQ ID NO: 1 or 1186 of SEQ ID NO: 3. The presence of the reference nucleotide at one or more of these positions indicates that the individual has a lower likelihood of having a vascular disease than an individual having the variant nucleotide at one or more of these positions, or a lower likelihood of having severe symptomology. In a particular embodiment, the individual is an individual at risk for development of a vascular disease.

The invention further relates to a method of diagnosing or aiding in the diagnosis of a disorder associated with the presence of one or more of (a) a serine at amino acid position 700 of SEO ID NO: 2; or (b) a proline at amino acid position 387 of SEQ ID

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NO: 4 in an individual. The method comprises obtaining a biological sample containing the TSP-1 and/or TSP-4 protein or relevant portion thereof from the individual and determining the amino acid present at one or more of the indicated amino acid positions, wherein presence of one or more of (a) a serine at amino acid position 700 of SEQ ID NO: 2; or (b) a proline at amino acid position 387 of SEQ ID NO: 4 is indicative of increased likelihood of said disorder in the individual as compared with an appropriate control, *e.g.*, an individual having the reference amino acid at one or more of said positions.

The invention further relates to a method of diagnosing or aiding in the diagnosis of a disorder associated with one or more of (a) a serine at amino acid position 700 of SEQ ID NO: 2; or (b) a proline at amino acid position 387 of SEQ ID NO: 4 in an individual. The method comprises obtaining a biological sample containing the TSP-1 and/or TSP-4 protein or relevant portion thereof from the individual and determining the amino acid present at one or more of the indicated amino acid positions, wherein presence of one or more of (a) an asparagine at amino acid position 700 of SEQ ID NO: 2; or (b) an alanine at amino acid position 387 of SEQ ID NO: 4 is indicative of decreased likelihood of said disorder in the individual as compared with an appropriate control, *e.g.*, an individual having the variant amino acid at one or more of said positions.

In one embodiment, the invention relates to a method for predicting the likelihood that an individual will have a vascular disease (or aiding in the diagnosis of a vascular disease), comprising the steps of obtaining a biological sample comprising the TSP-1 and/or TSP-4 protein or relevant portion thereof from an individual to be assessed and determining the amino acid present at one or more of amino acid positions 700 of SEQ ID NO: 2 or 387 of SEQ ID NO: 4. The presence of the reference amino acid at one or more of these positions indicates that the individual has a lower likelihood of having a vascular disease than an individual having the variant amino acid at one or more of these positions, or a lower likelihood of having severe symptomology. In a particular

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embodiment, the individual is an individual at risk for development of a vascular disease.

In another embodiment, the invention relates to pharmaceutical compositions comprising a reference TSP-1 and/or TSP-4 gene or gene product, or active portion thereof, for use in the treatment of vascular diseases. The invention further relates to the use of agonists and antagonists of TSP-1 and TSP-4 activity for use in the treatment of vascular diseases. In a particular embodiment the vascular disease is selected from the group consisting of atherosclerosis, coronary heart or artery disease, MI, stroke, peripheral vascular diseases, venous thromboembolism and pulmonary embolism. In a preferred embodiment, the vascular disease is selected from the group consisting of CAD and MI.

BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A-1D show the reference nucleotide (SEQ ID NO: 1) and amino acid (SEQ ID NO: 2) sequences for TSP-1.

Figs. 2A-2C show the reference nucleotide (SEQ ID NO: 3) and amino acid (SEQ ID NO: 4) sequences for TSP-4.

Fig. 3 shows a table providing detailed information about the SNPs identified herein. Column one shows the internal polymorphism identifier. Column two shows the accession number for the reference sequence in the TIGR database

(http://www.tigr.org/tdb/hgi/searching/hgi_reports.html). Column three shows the nucleotide position for the SNP iste. Column four shows the gene in which the polymorphism was identified. Column five shows the polymorphic site and additional flanking sequence on each side of the polymorphism. Column six shows the type of mutation produced by the polymorphism. Columns seven and eight show the reference and alternate (variant) nucleotides, respectively, for the SNP. Columns nine and ten show the reference and alternate (variant) amino acids, respectively, encoded by the alleles of the gene.

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DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to a gene which comprises a single nucleotide polymorphism (SNP) at a specific location. The gene which includes the SNP has at least two alleles, referred to herein as the reference allele and the variant allele. The reference allele (prototypical or wild type allele) has been designated arbitrarily and typically corresponds to the nucleotide sequence of the gene which has been deposited with GenBank or TIGR under a given Accession number. The variant allele differs from the reference allele by one nucleotide at the site(s) identified in the Table. The present invention also relates to variant alleles of the described genes and to complements of the variant alleles. The invention also relates to nucleic acid molecules which hybridize to and/or share identity with the variant alleles identified herein (or their complements) and which also comprise the variant nucleotide at the SNP site.

The invention further relates to portions of the variant alleles and portions of complements of the variant alleles which comprise (encompass) the site of the SNP and are at least 5 nucleotides in length. Portions can be, for example, 5-10, 5-15, 10-20, 5-25, 10-30, 10-50 or 10-100 bases long. For example, a portion of a variant allele which is 21 nucleotides in length includes the single nucleotide polymorphism (the nucleotide which differs from the reference allele at that site) and twenty additional nucleotides which flank the site in the variant allele. These nucleotides can be on one or both sides of the polymorphism. Polymorphisms which are the subject of this invention are defined in the Table with respect to the reference sequence deposited in GenBank or TIGR under the Accession number indicated. For example, the invention relates to a portion of a gene (e.g., AT3) having a nucleotide sequence as deposited in GenBank (e.g., U11270) comprising a single nucleotide polymorphism at a specific position (e.g., nucleotide 11918). The reference nucleotide for AT3 is shown in column 8, and the variant nucleotide is shown in column 9 of the Table. The nucleotide sequences of the invention can be double- or single-stranded.

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The invention further provides allele-specific oligonucleotides that hybridize to the reference or variant allele of a gene comprising a single nucleotide polymorphism or to the complement thereof. These oligonucleotides can be probes or primers.

The invention further provides a method of analyzing a nucleic acid from an individual. The method determines which base is present at any one of the polymorphic sites shown in the Table and/or Fig. 3. Optionally, a set of bases occupying a set of the polymorphic sites shown in the Table and/or Fig. 3 is determined. This type of analysis can be performed on a number of individuals, who are tested for the presence of a disease phenotype. The presence or absence of disease phenotype is then correlated with a base or set of bases present at the polymorphic site or sites in the individuals tested.

Thus, the invention further relates to a method of predicting the presence, absence, likelihood of the presence or absence, or severity of a particular phenotype or disorder associated with a particular genotype. The method comprises obtaining a nucleic acid sample from an individual and determining the identity of one or more bases (nucleotides) at polymorphic sites of genes described herein, wherein the presence of a particular base is correlated with a specified phenotype or disorder, thereby predicting the presence, absence, likelihood of the presence or absence, or severity of the phenotype or disorder in the individual.

20 DEFINITIONS

A nucleic acid molecule or oligonucleotide can be DNA or RNA, and single- or double-stranded. Nucleic acid molecules and oligonucleotides can be naturally occurring or synthetic, but are typically prepared by synthetic means. Preferred nucleic acid molecules and oligonucleotides of the invention include segments of DNA, or their complements, which include any one of the polymorphic sites shown in the Table. The segments can be between 5 and 250 bases, and, in specific embodiments, are between 5-10, 5-20, 10-20, 10-50, 20-50 or 10-100 bases. For example, the segment can be 21

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bases. The polymorphic site can occur within any position of the segment. The segments can be from any of the allelic forms of DNA shown in the Table.

As used herein, the terms "nucleotide", "base" and "nucleic acid" are intended to be equivalent. The terms "nucleotide sequence", "nucleic acid sequence", "nucleic acid molecule" and "segment" are intended to be equivalent.

Hybridization probes are oligonucleotides which bind in a base-specific manner to a complementary strand of nucleic acid. Such probes include peptide nucleic acids, as described in Nielsen *et al.*, *Science* 254, 1497-1500 (1991). Probes can be any length suitable for specific hybridization to the target nucleic acid sequence. The most appropriate length of the probe may vary depending upon the hybridization method in which it is being used; for example, particular lengths may be more appropriate for use in microfabricated arrays, while other lengths may be more suitable for use in classical hybridization methods. Such optimizations are known to the skilled artisan. Suitable probes and primers can range from about 5 nucleotides to about 30 nucleotides in length. For example, probes and primers can be 5, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 25, 26, 28 or 30 nucleotides in length. The probe or primer preferably overlaps at least one polymorphic site occupied by any of the possible variant nucleotides. The nucleotide sequence can correspond to the coding sequence of the allele or to the complement of the coding sequence of the allele.

As used herein, the term "primer" refers to a single-stranded oligonucleotide which acts as a point of initiation of template-directed DNA synthesis under appropriate conditions (e.g., in the presence of four different nucleoside triphosphates and an agent for polymerization, such as DNA or RNA polymerase or reverse transcriptase) in an appropriate buffer and at a suitable temperature. The appropriate length of a primer depends on the intended use of the primer, but typically ranges from 15 to 30 nucleotides. Short primer molecules generally require cooler temperatures to form sufficiently stable hybrid complexes with the template. A primer need not reflect the exact sequence of the template, but must be sufficiently complementary to hybridize with a template. The term primer site refers to the area of the target DNA to which a

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primer hybridizes. The term primer pair refers to a set of primers including a 5' (upstream) primer that hybridizes with the 5' end of the DNA sequence to be amplified and a 3' (downstream) primer that hybridizes with the complement of the 3' end of the sequence to be amplified.

As used herein, linkage describes the tendency of genes, alleles, loci or genetic markers to be inherited together as a result of their location on the same chromosome. It can be measured by percent recombination between the two genes, alleles, loci or genetic markers.

As used herein, polymorphism refers to the occurrence of two or more genetically determined alternative sequences or alleles in a population. A polymorphic marker or site is the locus at which divergence occurs. Preferred markers have at least two alleles, each occurring at frequency of greater than 1%, and more preferably greater than 10% or 20% of a selected population. A polymorphic locus may be as small as one base pair. Polymorphic markers include restriction fragment length polymorphisms, variable number of tandem repeats (VNTR's), hypervariable regions, minisatellites, dinucleotide repeats, trinucleotide repeats, tetranucleotide repeats, simple sequence repeats, and insertion elements such as Alu. The first identified allelic form is arbitrarily designated as the reference form and other allelic forms are designated as alternative or variant alleles. The allelic form occurring most frequently in a selected population is sometimes referred to as the wildtype form. Diploid organisms may be homozygous or heterozygous for allelic forms. A diallelic or biallelic polymorphism has two forms. A triallelic polymorphism has three forms.

Work described herein pertains to the resequencing of large numbers of genes in a large number of individuals to identify polymorphisms which can predispose individuals to disease. For example, polymorphisms in genes which are expressed in liver may predispose individuals to disorders of the liver. By altering amino acid sequence, SNPs may alter the function of the encoded proteins. The discovery of the SNP facilitates biochemical analysis of the variants and the development of assays to characterize the variants and to screen for pharmaceutical that would interact directly

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with on or another form of the protein. SNPs (including silent SNPs) also enable the development of specific DNA, RNA, or protein-based diagnostics that detect the presence or absence of the polymorphism in particular conditions.

A single nucleotide polymorphism occurs at a polymorphic site occupied by a single nucleotide, which is the site of variation between allelic sequences. The site is usually preceded by and followed by highly conserved sequences of the allele (e.g., sequences that vary in less than 1/100 or 1/1000 members of the populations).

A single nucleotide polymorphism usually arises due to substitution of one nucleotide for another at the polymorphic site. A transition is the replacement of one purine by another purine or one pyrimidine by another pyrimidine. A transversion is the replacement of a purine by a pyrimidine or vice versa. Single nucleotide polymorphisms can also arise from a deletion of a nucleotide or an insertion of a nucleotide relative to a reference allele. Typically the polymorphic site is occupied by a base other than the reference base. For example, where the reference allele contains the base "T" at the polymorphic site, the altered allele can contain a "C", "G" or "A" at the polymorphic site.

The invention also relates to nucleic acid molecules which hybridize to the variant alleles identified herein (or their complements) and which also comprise the variant nucleotide at the SNP site. Hybridizations are usually performed under stringent conditions, for example, at a salt concentration of no more than 1 M and a temperature of at least 25°C. For example, conditions of 5X SSPE (750 mM NaCl, 50 mM NaPhosphate, 5 mM EDTA, pH 7.4) and a temperature of 25-30°C, or equivalent conditions, are suitable for allele-specific probe hybridizations. Equivalent conditions can be determined by varying one or more of the parameters given as an example, as known in the art, while maintaining a similar degree of identity or similarity between the target nucleotide sequence and the primer or probe used.

The invention also relates to nucleic acid molecules which share substantial sequence identity to the variant alleles identified herein (or their complements) and which also comprise the variant nucleotide at the SNP site. Particularly preferred are

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nucleic acid molecules and fragments which have at least about 60%, preferably at least about 70, 80 or 85%, more preferably at least about 90%, even more preferably at least about 95%, and most preferably at least about 98% identity with nucleic acid molecules described herein. The percent identity of two nucleotide or amino acid sequences can be determined by aligning the sequences for optimal comparison purposes (e.g., gaps can be introduced in the sequence of a first sequence). The nucleotides or amino acids at corresponding positions are then compared, and the percent identity between the two sequences is a function of the number of identical positions shared by the sequences (i.e., % identity = # of identical positions/total # of positions x 100). In certain embodiments, the length of a sequence aligned for comparison purposes is at least 30%, preferably at least 40%, more preferably at least 60%, and even more preferably at least 70%, 80% or 90% of the length of the reference sequence. The actual comparison of the two sequences can be accomplished by well-known methods, for example, using a mathematical algorithm. A preferred, non-limiting example of such a mathematical algorithm is described in Karlin et al., Proc. Natl. Acad. Sci. USA, 90:5873-5877 (1993). Such an algorithm is incorporated into the NBLAST and XBLAST programs (version 2.0) as described in Altschul et al., Nucleic Acids Res., 25:389-3402 (1997). When utilizing BLAST and Gapped BLAST programs, the default parameters of the respective programs (e.g., NBLAST) can be used. See http://www.ncbi.nlm.nih.gov. In one embodiment, parameters for sequence comparison can be set at score=100, wordlength=12, or can be varied (e.g., W=5 or W=20).

The term "isolated" is used herein to indicate that the material in question exists in a physical milieu distinct from that in which it occurs in nature. For example, an isolated nucleic acid of the invention may be substantially isolated with respect to the complex cellular milieu in which it naturally occurs. In some instances, the isolated material will form part of a composition (for example, a crude extract containing other substances), buffer system or reagent mix. In other circumstance, the material may be purified to essential homogeneity, for example as determined by PAGE or column

chromatography such as HPLC. Preferably, an isolated nucleic acid comprises at least about 50, 80 or 90 percent (on a molar basis) of all macromolecular species present.

I. Novel Polymorphisms of the Invention

Some of the novel polymorphisms of the invention are shown in the Table.

- Columns one and two show designations for the indicated polymorphism. Column three shows the Genbank or TIGR Accession number for the wild type (or reference) allele. Column four shows the location of the polymorphic site in the nucleic acid sequence with reference to the Genbank or TIGR sequence shown in column three. Column five shows common names for the gene in which the polymorphism is located.
- 10 Column six shows the polymorphism and a portion of the 3' and 5' flanking sequence of the gene. Column seven shows the type of mutation; N, non-sense, S, silent, M, missense. Columns eight and nine show the reference and alternate nucleotides, respectively, at the polymorphic site. Columns ten and eleven show the reference and alternate amino acids, respectively, encoded by the reference and variant, respectively, alleles. Other novel polymorphisms of the invention are shown in Fig. 3.

II. Analysis of Polymorphisms

A. Preparation of Samples

Polymorphisms are detected in a target nucleic acid from an individual being analyzed. For assay of genomic DNA, virtually any biological sample (other than pure red blood cells) is suitable. For example, convenient tissue samples include whole blood, semen, saliva, tears, urine, fecal material, sweat, buccal, skin and hair. For assay of cDNA or mRNA, the tissue sample must be obtained from an organ in which the target nucleic acid is expressed. For example, if the target nucleic acid is a cytochrome P450, the liver is a suitable source.

25 Many of the methods described below require amplification of DNA from target samples. This can be accomplished by e.g., PCR. See generally PCR Technology: Principles and Applications for DNA Amplification (ed. H.A. Erlich, Freeman Press,

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NY, NY, 1992); PCR Protocols: A Guide to Methods and Applications (eds. Innis, et al., Academic Press, San Diego, CA, 1990); Mattila et al., Nucleic Acids Res. 19, 4967 (1991); Eckert et al., PCR Methods and Applications 1, 17 (1991); PCR (eds. McPherson et al., IRL Press, Oxford); and U.S. Patent 4,683,202.

Other suitable amplification methods include the ligase chain reaction (LCR) (see Wu and Wallace, *Genomics* 4, 560 (1989), Landegren *et al.*, *Science* 241, 1077 (1988), transcription amplification (Kwoh *et al.*, *Proc. Natl. Acad. Sci. USA* 86, 1173 (1989)), and self-sustained sequence replication (Guatelli *et al.*, *Proc. Nat. Acad. Sci. USA*, 87, 1874 (1990)) and nucleic acid based sequence amplification (NASBA). The latter two amplification methods involve isothermal reactions based on isothermal transcription, which produce both single stranded RNA (ssRNA) and double stranded DNA (dsDNA) as the amplification products in a ratio of about 30 or 100 to 1, respectively.

B. Detection of Polymorphisms in Target DNA

There are two distinct types of analysis of target DNA for detecting polymorphisms. The first type of analysis, sometimes referred to as de novo 15 characterization, is carried out to identify polymorphic sites not previously characterized (i.e., to identify new polymorphisms). This analysis compares target sequences in different individuals to identify points of variation, i.e., polymorphic sites. By analyzing groups of individuals representing the greatest ethnic diversity among humans 20 and greatest breed and species variety in plants and animals, patterns characteristic of the most common alleles/haplotypes of the locus can be identified, and the frequencies of such alleles/haplotypes in the population can be determined. Additional allelic frequencies can be determined for subpopulations characterized by criteria such as geography, race, or gender. The de novo identification of polymorphisms of the invention is described in the Examples section. The second type of analysis determines 25 which form(s) of a characterized (known) polymorphism are present in individuals under test. There are a variety of suitable procedures, which are discussed in turn.

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1. Allele-Specific Probes

The design and use of allele-specific probes for analyzing polymorphisms is described by e.g., Saiki *et al.*, *Nature* 324, 163-166 (1986); Dattagupta, EP 235,726, Saiki, WO 89/11548. Allele-specific probes can be designed that hybridize to a segment of target DNA from one individual but do not hybridize to the corresponding segment from another individual due to the presence of different polymorphic forms in the respective segments from the two individuals. Hybridization conditions should be sufficiently stringent that there is a significant difference in hybridization intensity between alleles, and preferably an essentially binary response, whereby a probe hybridizes to only one of the alleles. Some probes are designed to hybridize to a segment of target DNA such that the polymorphic site aligns with a central position (e.g., in a 15-mer at the 7 position; in a 16-mer, at either the 8 or 9 position) of the probe. This design of probe achieves good discrimination in hybridization between different allelic forms.

Allele-specific probes are often used in pairs, one member of a pair showing a perfect match to a reference form of a target sequence and the other member showing a perfect match to a variant form. Several pairs of probes can then be immobilized on the same support for simultaneous analysis of multiple polymorphisms within the same target sequence.

20 2. Tiling Arrays

The polymorphisms can also be identified by hybridization to nucleic acid arrays, some examples of which are described in WO 95/11995. One form of such arrays is described in the Examples section in connection with de novo identification of polymorphisms. The same array or a different array can be used for analysis of characterized polymorphisms. WO 95/11995 also describes subarrays that are optimized for detection of a variant form of a precharacterized polymorphism. Such a subarray contains probes designed to be complementary to a second reference sequence, which is an allelic variant of the first reference sequence. The second group of probes is

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designed by the same principles as described in the Examples, except that the probes exhibit complementarity to the second reference sequence. The inclusion of a second group (or further groups) can be particularly useful for analyzing short subsequences of the primary reference sequence in which multiple mutations are expected to occur within a short distance commensurate with the length of the probes (e.g., two or more mutations within 9 to 21 bases).

3. Allele-Specific Primers

An allele-specific primer hybridizes to a site on target DNA overlapping a polymorphism and only primes amplification of an allelic form to which the primer exhibits perfect complementarity. See Gibbs, *Nucleic Acid Res.* 17, 2427-2448 (1989). This primer is used in conjunction with a second primer which hybridizes at a distal site. Amplification proceeds from the two primers, resulting in a detectable product which indicates the particular allelic form is present. A control is usually performed with a second pair of primers, one of which shows a single base mismatch at the polymorphic site and the other of which exhibits perfect complementarity to a distal site. The single-base mismatch prevents amplification and no detectable product is formed. The method works best when the mismatch is included in the 3'-most position of the oligonucleotide aligned with the polymorphism because this position is most destabilizing to elongation from the primer (see, e.g., WO 93/22456).

20 4. Direct-Sequencing

The direct analysis of the sequence of polymorphisms of the present invention can be accomplished using either the dideoxy chain termination method or the Maxam - Gilbert method (see Sambrook *et al.*, *Molecular Cloning, A Laboratory Manual* (2nd Ed., CSHP, New York 1989); Zyskind *et al.*, *Recombinant DNA Laboratory Manual*, (Acad. Press, 1988)).

5. Denaturing Gradient Gel Electrophoresis

Amplification products generated using the polymerase chain reaction can be analyzed by the use of denaturing gradient gel electrophoresis. Different alleles can be identified based on the different sequence-dependent melting properties and electrophoretic migration of DNA in solution. Erlich, ed., *PCR Technology, Principles and Applications for DNA Amplification*, (W.H. Freeman and Co, New York, 1992), Chapter 7.

6. Single-Strand Conformation Polymorphism Analysis

Alleles of target sequences can be differentiated using single-strand conformation polymorphism analysis, which identifies base differences by alteration in electrophoretic migration of single stranded PCR products, as described in Orita *et al.*, *Proc. Nat. Acad. Sci.* 86, 2766-2770 (1989). Amplified PCR products can be generated as described above, and heated or otherwise denatured, to form single stranded amplification products. Single-stranded nucleic acids may refold or form secondary structures which are partially dependent on the base sequence. The different electrophoretic mobilities of single-stranded amplification products can be related to base-sequence differences between alleles of target sequences.

7. Single-Base Extension

An alternative method for identifying and analyzing polymorphisms is based on single-base extension (SBE) of a fluorescently-labeled primer coupled with fluorescence resonance energy transfer (FRET) between the label of the added base and the label of the primer. Typically, the method, such as that described by Chen *et al.*, (*PNAS 94*:10756-61 (1997), incorporated herein by reference) uses a locus-specific oligonucleotide primer labeled on the 5' terminus with 5-carboxyfluorescein (FAM).

25 This labeled primer is designed so that the 3' end is immediately adjacent to the polymorphic site of interest. The labeled primer is hybridized to the locus, and single base extension of the labeled primer is performed with fluorescently labeled

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dideoxyribonucleotides (ddNTPs) in dye-terminator sequencing fashion, except that no deoxyribonucleotides are present. An increase in fluorescence of the added ddNTP in response to excitation at the wavelength of the labeled primer is used to infer the identity of the added nucleotide.

5 III. Methods of Use

After determining polymorphic form(s) present in an individual at one or more polymorphic sites, this information can be used in a number of methods.

A. Forensics

Determination of which polymorphic forms occupy a set of polymorphic sites in an individual identifies a set of polymorphic forms that distinguishes the individual. *See generally* National Research Council, *The Evaluation of Forensic DNA Evidence* (Eds. Pollard *et al.*, National Academy Press, DC, 1996). The more sites that are analyzed, the lower the probability that the set of polymorphic forms in one individual is the same as that in an unrelated individual. Preferably, if multiple sites are analyzed, the sites are unlinked. Thus, polymorphisms of the invention are often used in conjunction with polymorphisms in distal genes. Preferred polymorphisms for use in forensics are biallelic because the population frequencies of two polymorphic forms can usually be determined with greater accuracy than those of multiple polymorphic forms at multi-allelic loci.

The capacity to identify a distinguishing or unique set of forensic markers in an individual is useful for forensic analysis. For example, one can determine whether a blood sample from a suspect matches a blood or other tissue sample from a crime scene by determining whether the set of polymorphic forms occupying selected polymorphic sites is the same in the suspect and the sample. If the set of polymorphic markers does not match between a suspect and a sample, it can be concluded (barring experimental error) that the suspect was not the source of the sample. If the set of markers does match, one can conclude that the DNA from the suspect is consistent with that found at

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the crime scene. If frequencies of the polymorphic forms at the loci tested have been determined (e.g., by analysis of a suitable population of individuals), one can perform a statistical analysis to determine the probability that a match of suspect and crime scene sample would occur by chance.

p(ID) is the probability that two random individuals have the same polymorphic or allelic form at a given polymorphic site. In biallelic loci, four genotypes are possible: AA, AB, BA, and BB. If alleles A and B occur in a haploid genome of the organism with frequencies x and y, the probability of each genotype in a diploid organism is (see WO 95/12607):

Homozygote: $p(AA) = x^2$

Homozygote: $p(BB)=y^2=(1-x)^2$

Single Heterozygote: p(AB)=p(BA)=xy=x(1-x)

Both Heterozygotes: p(AB+BA)= 2xy = 2x(1-x)

The probability of identity at one locus (i.e, the probability that two individuals, picked at random from a population will have identical polymorphic forms at a given locus) is given by the equation:

$$p(ID) = (x^2)^2 + (2xy)^2 + (y^2)^2$$
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These calculations can be extended for any number of polymorphic forms at a given locus. For example, the probability of identity p(ID) for a 3-allele system where the alleles have the frequencies in the population of x, y and z, respectively, is equal to the sum of the squares of the genotype frequencies:

$$p(ID) = x^4 + (2xy)^2 + (2yz)^2 + (2xz)^2 + z^4 + y^4$$

In a locus of n alleles, the appropriate binomial expansion is used to calculate p(ID) and p(exc).

The cumulative probability of identity (cum p(ID)) for each of multiple unlinked loci is determined by multiplying the probabilities provided by each locus.

$$cum p(ID) = p(ID1)p(ID2)p(ID3).... p(IDn)$$

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The cumulative probability of non-identity for n loci (i.e. the probability that two random individuals will be different at 1 or more loci) is given by the equation:

cum p(nonID) = 1-cum p(ID).

If several polymorphic loci are tested, the cumulative probability of non-identity for random individuals becomes very high (e.g., one billion to one). Such probabilities can be taken into account together with other evidence in determining the guilt or innocence of the suspect.

B. Paternity Testing

The object of paternity testing is usually to determine whether a male is the father of a child. In most cases, the mother of the child is known and thus, the mother's contribution to the child's genotype can be traced. Paternity testing investigates whether the part of the child's genotype not attributable to the mother is consistent with that of the putative father. Paternity testing can be performed by analyzing sets of polymorphisms in the putative father and the child.

If the set of polymorphisms in the child attributable to the father does not match the set of polymorphisms of the putative father, it can be concluded, barring experimental error, that the putative father is not the real father. If the set of polymorphisms in the child attributable to the father does match the set of polymorphisms of the putative father, a statistical calculation can be performed to determine the probability of coincidental match.

The probability of parentage exclusion (representing the probability that a random male will have a polymorphic form at a given polymorphic site that makes him incompatible as the father) is given by the equation (see WO 95/12607):

$$p(exc) = xy(1-xy)$$

where x and y are the population frequencies of alleles A and B of a biallelic polymorphic site.

(At a triallelic site p(exc) = xy(1-xy) + yz(1-yz) + xz(1-xz) + 3xyz(1-xyz)), where x, y and z and the respective population frequencies of alleles A, B and C).

The probability of non-exclusion is

p(non-exc) = 1 - p(exc)

The cumulative probability of non-exclusion (representing the value obtained when n loci are used) is thus:

5 cum p(non-exc) = p(non-exc1)p(non-exc2)p(non-exc3).... p(non-excn)

The cumulative probability of exclusion for n loci (representing the probability that a random male will be excluded)

cum p(exc) = 1 - cum p(non-exc).

If several polymorphic loci are included in the analysis, the cumulative
probability of exclusion of a random male is very high. This probability can be taken
into account in assessing the liability of a putative father whose polymorphic marker set
matches the child's polymorphic marker set attributable to his/her father.

C. Correlation of Polymorphisms with Phenotypic Traits

organism in different ways. Some polymorphisms occur within a protein coding sequence and contribute to phenotype by affecting protein structure. The effect may be neutral, beneficial or detrimental, or both beneficial and detrimental, depending on the circumstances. For example, a heterozygous sickle cell mutation confers resistance to malaria, but a homozygous sickle cell mutation is usually lethal. Other polymorphisms occur in noncoding regions but may exert phenotypic effects indirectly via influence on replication, transcription, and translation. A single polymorphism may affect more than one phenotypic trait. Likewise, a single phenotypic trait may be affected by polymorphisms in different genes. Further, some polymorphisms predispose an individual to a distinct mutation that is causally related to a certain phenotype.

25 Phenotypic traits include diseases that have known but hitherto unmapped genetic components (e.g., agammaglobulimenia, diabetes insipidus, Lesch-Nyhan syndrome, muscular dystrophy, Wiskott-Aldrich syndrome, Fabry's disease, familial hypercholesterolemia, polycystic kidney disease, hereditary spherocytosis, von

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Willebrand's disease, tuberous sclerosis, hereditary hemorrhagic telangicctasia, familial colonic polyposis, Ehlers-Danlos syndrome, osteogenesis imperfecta, and acute intermittent porphyria). Phenotypic traits also include symptoms of, or susceptibility to, multifactorial diseases of which a component is or may be genetic, such as autoimmune diseases, inflammation, cancer, diseases of the nervous system, and infection by pathogenic microorganisms. Some examples of autoimmune diseases include rheumatoid arthritis, multiple sclerosis, diabetes (insulin-dependent and non-independent), systemic lupus erythematosus and Graves disease. Some examples of cancers include cancers of the bladder, brain, breast, colon, esophagus, kidney, leukemia, liver, lung, oral cavity, ovary, pancreas, prostate, skin, stomach and uterus. Phenotypic traits also include characteristics such as longevity, appearance (e.g., baldness, obesity), strength, speed, endurance, fertility, and susceptibility or receptivity

The correlation of one or more polymorphisms with phenotypic traits can be facilitated by knowledge of the gene product of the wild type (reference) gene. The genes in which cSNPs of the present invention have been identified are genes which have been previously sequenced and characterized in one of their allelic forms.

to particular drugs or therapeutic treatments.

Correlation is performed for a population of individuals who have been tested for the presence or absence of a phenotypic trait of interest and for polymorphic markers sets. To perform such analysis, the presence or absence of a set of polymorphisms (i.e. a polymorphic set) is determined for a set of the individuals, some of whom exhibit a particular trait, and some of which exhibit lack of the trait. The alleles of each polymorphism of the set are then reviewed to determine whether the presence or absence of a particular allele is associated with the trait of interest. Correlation can be performed by standard statistical methods such as a K-squared test and statistically significant correlations between polymorphic form(s) and phenotypic characteristics are noted. For example, it might be found that the presence of allele A1 at polymorphism A correlates with heart disease. As a further example, it might be found that the combined

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presence of allele A1 at polymorphism A and allele B1 at polymorphism B correlates with increased milk production of a farm animal.

Such correlations can be exploited in several ways. In the case of a strong correlation between a set of one or more polymorphic forms and a disease for which treatment is available, detection of the polymorphic form set in a human or animal patient may justify immediate administration of treatment, or at least the institution of regular monitoring of the patient. Detection of a polymorphic form correlated with serious disease in a couple contemplating a family may also be valuable to the couple in their reproductive decisions. For example, the female partner might elect to undergo in vitro fertilization to avoid the possibility of transmitting such a polymorphism from her husband to her offspring. In the case of a weaker, but still statistically significant correlation between a polymorphic set and human disease, immediate therapeutic intervention or monitoring may not be justified. Nevertheless, the patient can be motivated to begin simple life-style changes (e.g., diet, exercise) that can be accomplished at little cost to the patient but confer potential benefits in reducing the risk of conditions to which the patient may have increased susceptibility by virtue of variant alleles. Identification of a polymorphic set in a patient correlated with enhanced receptiveness to one of several treatment regimes for a disease indicates that this treatment regime should be followed.

For animals and plants, correlations between characteristics and phenotype are useful for breeding for desired characteristics. For example, Beitz *et al.*, US 5,292,639 discuss use of bovine mitochondrial polymorphisms in a breeding program to improve milk production in cows. To evaluate the effect of mtDNA D-loop sequence polymorphism on milk production, each cow was assigned a value of 1 if variant or 0 if wildtype with respect to a prototypical mitochondrial DNA sequence at each of 17 locations considered. Each production trait was analyzed individually with the following animal model:

$$Y_{ijkpn} = \mu + YS_i + P_j + X_k + \beta_1 + ... \beta_{17} + PE_n + a_n + e_p$$

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where Y_{ijknp} is the milk, fat, fat percentage, SNF, SNF percentage, energy concentration, or lactation energy record; μ is an overall mean; YS_i is the effect common to all cows calving in year-season; X_k is the effect common to cows in either the high or average selection line; β_1 to β_{17} are the binomial regressions of production record on mtDNA D-loop sequence polymorphisms; PE_n is permanent environmental effect common to all records of cow n; a_n is effect of animal n and is composed of the additive genetic contribution of sire and dam breeding values and a Mendelian sampling effect; and e_p is a random residual. It was found that eleven of seventeen polymorphisms tested influenced at least one production trait. Bovines having the best polymorphic forms for milk production at these eleven loci are used as parents for breeding the next generation of the herd.

D. Genetic Mapping of Phenotypic Traits

The previous section concerns identifying correlations between phenotypic traits and polymorphisms that directly or indirectly contribute to those traits. The present section describes identification of a physical linkage between a genetic locus associated with a trait of interest and polymorphic markers that are not associated with the trait, but are in physical proximity with the genetic locus responsible for the trait and cosegregate with it. Such analysis is useful for mapping a genetic locus associated with a phenotypic trait to a chromosomal position, and thereby cloning gene(s) responsible for the trait. See Lander et al., Proc. Natl. Acad. Sci. (USA) 83, 7353-7357 (1986); Lander et al., Proc. Natl. Acad. Sci. (USA) 84, 2363-2367 (1987); Donis-Keller et al., Cell 51, 319-337 (1987); Lander et al., Genetics 121, 185-199 (1989)). Genes localized by linkage can be cloned by a process known as directional cloning. See Wainwright, Med. J. Australia 159, 170-174 (1993); Collins, Nature Genetics 1, 3-6 (1992).

Linkage studies are typically performed on members of a family. Available members of the family are characterized for the presence or absence of a phenotypic trait and for a set of polymorphic markers. The distribution of polymorphic markers in an informative meiosis is then analyzed to determine which polymorphic markers co-

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segregate with a phenotypic trait. *See, e.g.*, Kerem *et al.*, *Science* 245, 1073-1080 (1989); Monaco *et al.*, *Nature* 316, 842 (1985); Yamoka *et al.*, *Neurology* 40, 222-226 (1990); Rossiter *et al.*, *FASEB Journal* 5, 21-27 (1991).

Linkage is analyzed by calculation of LOD (log of the odds) values. A lod value is the relative likelihood of obtaining observed segregation data for a marker and a genetic locus when the two are located at a recombination fraction θ , versus the situation in which the two are not linked, and thus segregating independently (Thompson & Thompson, Genetics in Medicine (5th ed, W.B. Saunders Company, Philadelphia, 1991); Strachan, "Mapping the human genome" in *The Human Genome* (BIOS Scientific Publishers Ltd, Oxford), Chapter 4). A series of likelihood ratios are calculated at various recombination fractions (θ), ranging from $\theta = 0.0$ (coincident loci) to $\theta = 0.50$ (unlinked). Thus, the likelihood at a given value of θ is: probability of data if loci linked at θ to probability of data if loci unlinked. The computed likelihoods are usually expressed as the log₁₀ of this ratio (i.e., a lod score). For example, a lod score of 3 indicates 1000:1 odds against an apparent observed linkage being a coincidence. The use of logarithms allows data collected from different families to be combined by simple addition. Computer programs are available for the calculation of lod scores for differing values of θ (e.g., LIPED, MLINK (Lathrop, Proc. Nat. Acad. Sci. (USA) 81, 3443-3446 (1984)). For any particular lod score, a recombination fraction may be determined from mathematical tables. See Smith et al., Mathematical tables for research workers in human genetics (Churchill, London, 1961); Smith, Ann. Hum. Genet. 32, 127-150 (1968). The value of θ at which the lod score is the highest is considered to be the best estimate of the recombination fraction.

Positive lod score values suggest that the two loci are linked, whereas negative values suggest that linkage is less likely (at that value of θ) than the possibility that the two loci are unlinked. By convention, a combined lod score of +3 or greater (equivalent to greater than 1000:1 odds in favor of linkage) is considered definitive evidence that two loci are linked. Similarly, by convention, a negative lod score of -2 or less is taken as definitive evidence against linkage of the two loci being compared. Negative linkage

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data are useful in excluding a chromosome or a segment thereof from consideration. The search focuses on the remaining non-excluded chromosomal locations.

IV. Modified Polypeptides and Gene Sequences

The invention further provides variant forms of nucleic acids and corresponding proteins. The nucleic acids comprise one of the sequences described in the Table, column 5, in which the polymorphic position is occupied by one of the alternative bases for that position. Some nucleic acids encode full-length variant forms of proteins. Similarly, variant proteins have the prototypical amino acid sequences encoded by nucleic acid sequences shown in the Table, column 5, (read so as to be in-frame with the full-length coding sequence of which it is a component) except at an amino acid encoded by a codon including one of the polymorphic positions shown in the Table. That position is occupied by the amino acid coded by the corresponding codon in any of the alternative forms shown in the Table.

Variant genes can be expressed in an expression vector in which a variant gene is operably linked to a native or other promoter. Usually, the promoter is a eukaryotic promoter for expression in a mammalian cell. The transcription regulation sequences typically include a heterologous promoter and optionally an enhancer which is recognized by the host. The selection of an appropriate promoter, for example trp, lac, phage promoters, glycolytic enzyme promoters and tRNA promoters, depends on the host selected. Commercially available expression vectors can be used. Vectors can include host-recognized replication systems, amplifiable genes, selectable markers, host sequences useful for insertion into the host genome, and the like.

The means of introducing the expression construct into a host cell varies depending upon the particular construction and the target host. Suitable means include fusion, conjugation, transfection, transduction, electroporation or injection, as described in Sambrook, *supra*. A wide variety of host cells can be employed for expression of the variant gene, both prokaryotic and eukaryotic. Suitable host cells include bacteria such as *E. coli*, yeast, filamentous fungi, insect cells, mammalian cells, typically

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immortalized, *e.g.*, mouse, CHO, human and monkey cell lines and derivatives thereof. Preferred host cells are able to process the variant gene product to produce an appropriate mature polypeptide. Processing includes glycosylation, ubiquitination, disulfide bond formation, general post-translational modification, and the like. As used herein, "gene product" includes mRNA, peptide and protein products.

The protein may be isolated by conventional means of protein biochemistry and purification to obtain a substantially pure product, *i.e.*, 80, 95 or 99% free of cell component contaminants, as described in Jacoby, *Methods in Enzymology* Volume 104, Academic Press, New York (1984); Scopes, *Protein Purification, Principles and Practice*, 2nd Edition, Springer-Verlag, New York (1987); and Deutscher (ed), *Guide to Protein Purification, Methods in Enzymology*, Vol. 182 (1990). If the protein is secreted, it can be isolated from the supernatant in which the host cell is grown. If not secreted, the protein can be isolated from a lysate of the host cells.

The invention further provides transgenic nonhuman animals capable of expressing an exogenous variant gene and/or having one or both alleles of an endogenous variant gene inactivated. Expression of an exogenous variant gene is usually achieved by operably linking the gene to a promoter and optionally an enhancer, and microinjecting the construct into a zygote. See Hogan et al., "Manipulating the Mouse Embryo, A Laboratory Manual," Cold Spring Harbor Laboratory. Inactivation of endogenous variant genes can be achieved by forming a transgene in which a cloned variant gene is inactivated by insertion of a positive selection marker. See Capecchi, Science 244, 1288-1292 (1989). The transgene is then introduced into an embryonic stem cell, where it undergoes homologous recombination with an endogenous variant gene. Mice and other rodents are preferred animals. Such animals provide useful drug screening systems.

In addition to substantially full-length polypeptides expressed by variant genes, the present invention includes biologically active fragments of the polypeptides, or analogs thereof, including organic molecules which simulate the interactions of the peptides. Biologically active fragments include any portion of the full-length

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polypeptide which confers a biological function on the variant gene product, including ligand binding, and antibody binding. Ligand binding includes binding by nucleic acids, proteins or polypeptides, small biologically active molecules, or large cellular structures.

Polyclonal and/or monoclonal antibodies that specifically bind to variant gene products but not to corresponding prototypical gene products are also provided. Antibodies can be made by injecting mice or other animals with the variant gene product or synthetic peptide fragments thereof. Monoclonal antibodies are screened as are described, for example, in Harlow & Lane, *Antibodies, A Laboratory Manual*, Cold Spring Harbor Press, New York (1988); Goding, *Monoclonal antibodies, Principles and Practice* (2d ed.) Academic Press, New York (1986). Monoclonal antibodies are tested for specific immunoreactivity with a variant gene product and lack of immunoreactivity to the corresponding prototypical gene product. These antibodies are useful in diagnostic assays for detection of the variant form, or as an active ingredient in a pharmaceutical composition.

V. Kits

The invention further provides kits comprising at least one allele-specific oligonucleotide as described herein. Often, the kits contain one or more pairs of allele-specific oligonucleotides hybridizing to different forms of a polymorphism. In some kits, the allele-specific oligonucleotides are provided immobilized to a substrate. For example, the same substrate can comprise allele-specific oligonucleotide probes for detecting at least 10, 100 or all of the polymorphisms shown in the Table. Optional additional components of the kit include, for example, restriction enzymes, reverse-transcriptase or polymerase, the substrate nucleoside triphosphates, means used to label (for example, an avidin-enzyme conjugate and enzyme substrate and chromogen if the label is biotin), and the appropriate buffers for reverse transcription, PCR, or hybridization reactions. Usually, the kit also contains instructions for carrying out the methods.

The thrombospondins are a family of extracellular matrix (ECM) glycoproteins that modulate many cell behaviors including adhesion, migration, and proliferation. Thrombospondins (also known as thrombin sensitive proteins or TSPs) are large molecular weight glycoproteins composed of three identical disulfide-linked polypeptide chains. TSPs are stored in the alpha-granules of platelets and secreted by a variety of mesenchymal and epithelial cells (Majack *et al.*, *Cell Membrane 3:57-77* (1987)). Platelets secrete TSPs when activated in the blood by such physiological agonists such as thrombin. TSPs have lectin properties and a broad function in the regulation of fibrinolysis and as a component of the ECM, and are one of a group of ECM proteins which have adhesive properties. TSPs bind to fibronectin and fibrinogen (Lahav *et al.*, *Eur J Biochem 145*:151-6 (1984)), and these proteins are known to be involved in platelet adhesion to substratum and platelet aggregation (Leung, *J Clin Invest 74*:1764-1772 (1986)).

Recent work has implicated TSPs in response of cells to growth factors. 15 Submitogenic doses of PDGF induce a rapid but transitory, increase in TSP synthesis and secretion by rat aortic smooth muscle cells (Majack et al., J Biol Chem 101:1059-70 (1985)). PDGF responsiveness to TSP synthesis in glial cells has also been shown (Asch et al., Proc Natl Acad Sci 83:2904-8 (1986)). TSP mRNA levels rise rapidly in response to PDGF (Majack et al., J Biol Chem 262:8821-5 (1987)). TSPs act 20 synergistically with epidermal growth factor to increase DNA synthesis in smooth muscle cells (Majack et al., Proc Natl Acad Sci 83:9050-4 (1986)), and monoclonal antibodies to TSPs inhibit smooth muscle cell proliferation (Majack et al., J Biol Chem 106:415-22 (1988)). TSPs modulate local adhesions in endothelial cells, and TSPs, particularly TSP-1 primarily derived from platelet granules, are known to be an important activator of transforming growth factor beta-1 (TGFB-1) (Crawford et al., Cell 93:1159 (1998)) and appear to be a potential link between platelet-thrombosis and development of atherosclerosis.

To determine pivotal genes associated with premature coronary artery disease, we analyzed DNA from 347 patients with MI or coronary revascularization before age 40

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(men) or 45 (women) and 422 general population controls. Cases were drawn (one per family) from a retrospective collection of sibling pairs with premature CAD. Controls were ascertained through random-digit dialing. Both cases and controls were Caucasian. A complete database of phenotypic and laboratory variables for the affected patients afforded logistic regression to control for age, diabetes, body mass index, gender.

Thrombospondin (TSP) 4 and 1 emerged as important SNPs associated with premature CAD and MI. For CAD, 148 of 347 patients carried at least one copy of the TSP-4 variant compared with 142 of 422 control subjects; adjusted odds ratio 1.47, p=0.01. For premature MI, the association was even stronger: 91 of 187 cases vs. 142 of 422 controls had the variant; adjusted odds ratio 2.08, p=0.0003. The TSP-1 SNP was rare. Nonetheless, homozygosity for the variant allele gave an adjusted odds ratio of 9.5, p=.04.

Specific reference nucleotide (SEQ ID NO: 1) and amino acid (SEQ ID NO: 2) sequences for TSP-1 are shown in Figs. 1A-1D. Specific reference nucleotide (SEQ ID NO: 3) and amino acid (SEQ ID NO: 4) sequences for TSP-4 are shown in Figs. 2A-2C. It is understood that the invention is not limited by these exemplified reference sequences, as variants of these sequences which differ at locations other than the SNP sites identified herein can also be utilized. The skilled artisan can readily determine the SNP sites in these other reference sequences which correspond to the SNP sites identified herein by aligning the sequence of interest with the reference sequences specifically disclosed herein, and programs for performing such alignments are commercially available. For example, the ALIGN program in the GCG software package can be used, utilizing a PAM120 weight residue table, a gap length penalty of 12 and a gap penalty of 4, for example.

Two SNPs have been specifically studied as described herein. The first (G334u4) is a change from A (reference nucleotide) to G (alternate or variant nucleotide) at nucleotide position 2210 of the nucleic acid sequence of TSP-1 (Figs. 1A-1D), resulting in a missense amino acid mutation from asparagine (reference) to serine (alternate) at

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amino acid 700. The second SNP (G355u2) is a change from G (reference) to C (alternate) at nucleotide position 1186 of the nucleic acid sequence of TSP-4 (Figs. 2A-2C), resulting in a missense amino acid alteration from alanine (reference) to proline (alternate) at amino acid 387. With respect to the G355u2 SNP, individuals with CAD carried at least one copy of the variant "C" allele more frequently than control individuals (43% as compared with 34%). With respect to the G355u2 SNP, individuals with MI carried at least one copy of the variant "C" allele more frequently than control individuals (49% as compared with 34%). With respect to the G334u4 SNP, individuals with CAD carried two copies of the variant "G" allele more frequently than control individuals (1.7% as compared with 0.2%). With respect to the G334u4 SNP, individuals with MI carried two copies of the variant "G" allele more frequently than control individuals (2% as compared with 0.2%).

As used herein, the term "polymorphism" refers to the occurrence of two or more genetically determined alternative sequences or alleles in a population. A polymorphic marker or site is the locus at which divergence occurs. Preferred markers have at least two alleles, each occurring at frequency of greater than 1%, and more preferably greater than 10% or 20% of a selected population. A polymorphic locus may be as small as one base pair, in which case it is referred to as a single nucleotide polymorphism (SNP).

Thus, the invention relates to a method for predicting the likelihood that an individual will have a vascular disease, or for aiding in the diagnosis of a vascular disease, or predicting the likelihood of having altered symptomology associated with a vascular disease, comprising the steps of obtaining a DNA sample from an individual to be assessed and determining the nucleotide present at one or more of nucleotide positions 2210 of the TSP-1 gene or 1186 of the TSP-4 gene. In a preferred embodiment, the nucleotides present at both of these nucleotide positions are determined. In one embodiment the TSP-1 gene has the nucleotide sequence of SEQ ID NO: 1 and the TSP-4 gene has the nucleotide sequence of SEQ ID NO: 3. The presence of one or more of a G (the variant nucleotide) at position 2210 of SEQ ID NO: 1 or a C (the variant nucleotide) at position 1186 of SEQ ID NO: 1186 indicates that the

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individual has a greater likelihood of having a vascular disease, or a greater likelihood of having severe symptomology associated with a vascular disease, than if that individual had the reference nucleotide at one or more of these positions. Conversely, the presence of one or more of an A (the reference nucleotide) at position 2210 of SEQ ID NO: 1 or a G (the reference nucleotide) at position 1186 of SEQ ID NO: 3 indicates that the individual has a reduced likelihood of having a vascular disease or a likelihood of having reduced symptomology associated with a vascular disease than if that individual had the variant nucleotide at one or more of these positions.

In a particular embodiment, the individual is an individual at risk for development of a vascular disease. In another embodiment the individual exhibits clinical symptomology associated with a vascular disease. In one embodiment, the individual has been clinically diagnosed as having a vascular disease. Vascular diseases include, but are not limited to, atherosclerosis, coronary heart disease, myocardial infarction (MI), stroke, peripheral vascular diseases, venous thromboembolism and pulmonary embolism. In preferred embodiments, the vascular disease is CAD or MI.

The genetic material to be assessed can be obtained from any nucleated cell from the individual. For assay of genomic DNA, virtually any biological sample (other than pure red blood cells) is suitable. For example, convenient tissue samples include whole blood, semen, saliva, tears, urine, fecal material, sweat, skin and hair. For assay of cDNA or mRNA, the tissue sample must be obtained from a tissue or organ in which the target nucleic acid is expressed.

Many of the methods described herein require amplification of DNA from target samples. This can be accomplished by e.g., PCR. See generally PCR Technology: Principles and Applications for DNA Amplification (ed. H.A. Erlich, Freeman Press, NY, NY, 1992); PCR Protocols: A Guide to Methods and Applications (eds. Innis, et al., Academic Press, San Diego, CA, 1990); Mattila et al., Nucleic Acids Res. 19, 4967 (1991); Eckert et al., PCR Methods and Applications 1, 17 (1991); PCR (eds. McPherson et al., IRL Press, Oxford); and U.S. Patent 4,683,202.

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Other suitable amplification methods include the ligase chain reaction (LCR) (see Wu and Wallace, *Genomics* 4, 560 (1989), Landegren *et al.*, *Science* 241, 1077 (1988), transcription amplification (Kwoh *et al.*, *Proc. Natl. Acad. Sci. USA* 86, 1173 (1989)), and self-sustained sequence replication (Guatelli *et al.*, *Proc. Nat. Acad. Sci. USA*, 87, 1874 (1990)) and nucleic acid based sequence amplification (NASBA). The latter two amplification methods involve isothermal reactions based on isothermal transcription, which produce both single stranded RNA (ssRNA) and double stranded DNA (dsDNA) as the amplification products in a ratio of about 30 or 100 to 1, respectively.

The nucleotide which occupies the polymorphic site of interest (e.g., nucleotide position 2210 in TSP-1 and/or nucleotide position 1186 in TSP-4) can be identified by a variety of methods, such as Southern analysis of genomic DNA; direct mutation analysis by restriction enzyme digestion; Northern analysis of RNA; denaturing high pressure liquid chromatography (DHPLC); gene isolation and sequencing; hybridization of an allele-specific oligonucleotide with amplified gene products; single base extension (SBE). In a preferred embodiment, determination of the allelic form of TSP is carried out using SBE-FRET methods as described herein, or using chip-based oligonucleotide arrays as described herein.

The invention also relates to a method for predicting the likelihood that an individual will have a vascular disease, or for aiding in the diagnosis of a vascular disease, or predicting the likelihood of having altered symptomology associated with a vascular disease, comprising the steps of obtaining a biological sample comprising TSP-1 and/or TSP-4 protein or relevant portion thereof from an individual to be assessed and determining the amino acid present at one or more of amino acid positions 700 of the TSP-1 gene product (e.g., as exemplified by SEQ ID NO: 2) or 387 of the TSP-4 gene product (e.g., as exemplified by SEQ ID NO: 4). In a preferred embodiment, the amino acids present at both of these amino acid positions are determined. As used herein, the term "relevant portion" of the TSP-1 and TSP-4 proteins is intended to encompass any portion of the protein which comprises the polymorphic amino acid positions. The presence of one or more of a serine (the variant amino acid) at position 700 of SEQ ID

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NO: 2, or a proline (the variant amino acid) at position 387 of SEQ ID NO: 4 indicates that the individual has a greater likelihood of having a vascular disease, or a greater likelihood of having severe symptomology associated with a vascular disease, than if that individual had the reference amino acid at one or more of these positions.

5 Conversely, the presence of one or more of an asparagine (the reference amino acid) at position 700 of SEQ ID NO: 2, or an alanine (the reference amino acid) at position 387 of SEQ I D NO: 4 indicates that the individual has a reduced likelihood of having a vascular disease or a likelihood of having reduced symptomology associated with a vascular disease, than if that individual had the varaint amino acid at one or more of these positions.

In a particular embodiment, the individual is an individual at risk for development of a vascular disease. In another embodiment the individual exhibits clinical symptomology associated with a vascular disease. In one embodiment, the individual has been clinically diagnosed as having a vascular disease.

In this embodiment of the invention, the biological sample contains protein molecules from the test subject. *In vitro* techniques for detection of protein include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations and immunofluorescence. Furthermore, *in vivo* techniques for detection of protein include introducing into a subject a labeled anti-protein antibody. For example, the antibody can be labeled with a radioactive marker whose presence and location in a subject can be detected by standard imaging techniques. Polyclonal and/or monoclonal antibodies that specifically bind to variant gene products but not to corresponding reference gene products, and vice versa, are also provided. Antibodies can be made by injecting mice or other animals with the variant gene product or synthetic peptide fragments thereof comprising the variant portion. Monoclonal antibodies are screened as are described, for example, in Harlow & Lane, *Antibodies, A Laboratory Manual*, Cold Spring Harbor Press, New York (1988); Goding, *Monoclonal antibodies, Principles and Practice* (2d ed.) Academic Press, New York (1986). Monoclonal antibodies are tested for specific immunoreactivity with a variant gene product and lack

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of immunoreactivity to the corresponding prototypical gene product. These antibodies are useful in diagnostic assays for detection of the variant form, or as an active ingredient in a pharmaceutical composition.

The polymorphisms of the invention may be associated with vascular disease in different ways. The polymorphisms may exert phenotypic effects indirectly via influence on replication, transcription, and translation. Additionally, the described polymorphisms may predispose an individual to a distinct mutation that is causally related to a certain phenotype, such as susceptibility or resistance to vascular disease and related disorders. The discovery of the polymorphisms and their correlation with CAD and MI facilitates biochemical analysis of the variant and reference forms and the development of assays to characterize the variant and reference forms and to screen for pharmaceutical agents that interact directly with one or another form of the protein.

Alternatively, these particular polymorphisms may belong to a group of two or more polymorphisms in the TSP gene(s) which contributes to the presence, absence or severity of vascular disease. An assessment of other polymorphisms within the TSP gene(s) can be undertaken, and the separate and combined effects of these polymorphisms, as well as alternations in other, distinct genes, on the vascular disease phenotype can be assessed.

Correlation between a particular phenotype, e.g., the CAD or MI phenotype, and the presence or absence of a particular allele is performed for a population of individuals who have been tested for the presence or absence of the phenotype. Correlation can be performed by standard statistical methods such as a Chi-squared test and statistically significant correlations between polymorphic form(s) and phenotypic characteristics are noted. This correlation can be exploited in several ways. In the case of a strong correlation between a particular polymorphic form, e.g., the variant allele for TSP-1 and/or TSP-4, and a disease for which treatment is available, detection of the polymorphic form in an individual may justify immediate administration of treatment, or at least the institution of regular monitoring of the individual. Detection of a polymorphic form correlated with a disorder in a couple contemplating a family may

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also be valuable to the couple in their reproductive decisions. For example, the female partner might elect to undergo *in vitro* fertilization to avoid the possibility of transmitting such a polymorphism from her husband to her offspring. In the case of a weaker, but still statistically significant correlation between a polymorphic form and a particular disorder, immediate therapeutic intervention or monitoring may not be justified. Nevertheless, the individual can be motivated to begin simple life-style changes (e.g., diet modification, therapy or counseling) that can be accomplished at little cost to the individual but confer potential benefits in reducing the risk of conditions to which the individual may have increased susceptibility by virtue of the particular allele. Furthermore, identification of a polymorphic form correlated with enhanced receptiveness to one of several treatment regimes for a disorder indicates that this treatment regimen should be followed for the individual in question.

Furthermore, it may be possible to identify a physical linkage between a genetic locus associated with a trait of interest (e.g., CAD or MI) and polymorphic markers that are or are not associated with the trait, but are in physical proximity with the genetic locus responsible for the trait and co-segregate with it. Such analysis is useful for mapping a genetic locus associated with a phenotypic trait to a chromosomal position, and thereby cloning gene(s) responsible for the trait. See Lander et al., Proc. Natl. Acad. Sci. (USA) 83, 7353-7357 (1986); Lander et al., Proc. Natl. Acad. Sci. (USA) 84, 2363-2367 (1987); Donis-Keller et al., Cell 51, 319-337 (1987); Lander et al., Genetics 121, 185-199 (1989)). Genes localized by linkage can be cloned by a process known as directional cloning. See Wainwright, Med. J. Australia 159, 170-174 (1993); Collins, Nature Genetics 1, 3-6 (1992). Linkage studies are discussed in more detail above.

In another embodiment, the invention relates to pharmaceutical compositions comprising a reference TSP-1 and/or TSP-4 gene or gene product for use in the treatment of vascular disease, e.g., CAD and MI. As used herein, a reference TSP gene product is intended to mean gene products which are encoded by the reference allele of the TSP gene. In addition to substantially full-length polypeptides expressed by the genes, the present invention includes biologically active fragments of the polypeptides,

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or analogs thereof, including organic molecules which simulate the interactions of the peptides. Biologically active fragments include any portion of the full-length polypeptide which confers a biological function on the variant gene product, including ligand binding, and antibody binding. Ligand binding includes binding by nucleic acids, proteins or polypeptides, small biologically active molecules, or large cellular structures.

For instance, the polypeptide or protein, or fragment thereof, of the present invention can be formulated with a physiologically acceptable medium to prepare a pharmaceutical composition. The particular physiological medium may include, but is not limited to, water, buffered saline, polyols (e.g., glycerol, propylene glycol, liquid polyethylene glycol) and dextrose solutions. The optimum concentration of the active ingredient(s) in the chosen medium can be determined empirically, according to procedures well known to medicinal chemists, and will depend on the ultimate pharmaceutical formulation desired. Methods of introduction of exogenous peptides at the site of treatment include, but are not limited to, intradermal, intramuscular, intraperitoneal, intravenous, subcutaneous, oral and intranasal. Other suitable methods of introduction can also include rechargeable or biodegradable devices and slow release polymeric devices. The pharmaceutical compositions of this invention can also be administered as part of a combinatorial therapy with other agents and treatment regimens.

The invention further pertains to compositions, e.g., vectors, comprising a nucleotide sequence encoding reference or variant TSP-1 and/or TSP-4 gene products. For example, reference genes can be expressed in an expression vector in which a reference gene is operably linked to a native or other promoter. Usually, the promoter is a eukaryotic promoter for expression in a mammalian cell. The transcription regulation sequences typically include a heterologous promoter and optionally an enhancer which is recognized by the host. The selection of an appropriate promoter, for example trp, lac, phage promoters, glycolytic enzyme promoters and tRNA promoters, depends on the host selected. Commercially available expression vectors can be used. Vectors can

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include host-recognized replication systems, amplifiable genes, selectable markers, host sequences useful for insertion into the host genome, and the like.

The means of introducing the expression construct into a host cell varies depending upon the particular construction and the target host. Suitable means include fusion, conjugation, transfection, transduction, electroporation or injection, as described in Sambrook, *supra*. A wide variety of host cells can be employed for expression of the variant gene, both prokaryotic and eukaryotic. Suitable host cells include bacteria such as *E. coli*, yeast, filamentous fungi, insect cells, mammalian cells, typically immortalized, *e.g.*, mouse, CHO, human and monkey cell lines and derivatives thereof. Preferred host cells are able to process the variant gene product to produce an appropriate mature polypeptide. Processing includes glycosylation, ubiquitination, disulfide bond formation, general post-translational modification, and the like.

It is also contemplated that cells can be engineered to express the reference allele of the invention by gene therapy methods. For example, DNA encoding the reference TSP gene product, or an active fragment or derivative thereof, can be introduced into an expression vector, such as a viral vector, and the vector can be introduced into appropriate cells in an animal. In such a method, the cell population can be engineered to inducibly or constitutively express active reference TSP gene product. In a preferred embodiment, the vector is delivered to the bone marrow, for example as described in Corey et al. (Science 244:1275-1281 (1989)).

The invention further relates to the use of compositions (i.e., agonists) which enhance or increase the activity of the reference (or variant) TSP (e.g., TSP-1 or TSP-4) gene product, or a functional portion thereof, for use in the treatment of vascular disease. The invention also relates to the use of compositions (i.e., antagonists) which reduce or decrease the activity of the variant (or reference) TSP (e.g., TSP-1 or TSP-4) gene product, or a functional portion thereof, for use in the treatment of vascular disease.

The invention also relates to constructs which comprise a vector into which a sequence of the invention has been inserted in a sense or antisense orientation. For

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example, a vector comprising a nucleotide sequence which is antisense to the variant TSP-1 or TSP-4 allele may be used as an antagonist of the activity of the TSP-1 or TSP-4 variant allele. Alternatively, a vector comprising a nucleotide sequence of the TSP-1 or TSP-4 reference allele may be used therapeutically to treat vascular diseases. As used herein, the term "vector" refers to a nucleic acid molecule capable of transporting another nucleic acid to which it has been linked. One type of vector is a "plasmid", which refers to a circular double stranded DNA loop into which additional DNA segments can be ligated. Another type of vector is a viral vector, wherein additional DNA segments can be ligated into the viral genome. Certain vectors are capable of autonomous replication in a host cell into which they are introduced (e.g., bacterial vectors having a bacterial origin of replication and episomal mammalian vectors). Other vectors (e.g., non-episomal mammalian vectors) are integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome. Moreover, certain vectors, expression vectors, are capable of directing the expression of genes to which they are operably linked. In general, expression vectors of utility in recombinant DNA techniques are often in the form of plasmids (vectors). However, the invention is intended to include such other forms of expression vectors, such as viral vectors (e.g., replication defective retroviruses, adenoviruses and adeno-associated viruses) that serve equivalent functions.

Preferred recombinant expression vectors of the invention comprise a nucleic acid of the invention in a form suitable for expression of the nucleic acid in a host cell. This means that the recombinant expression vectors include one or more regulatory sequences, selected on the basis of the host cells to be used for expression, which is operably linked to the nucleic acid sequence to be expressed. Within a recombinant expression vector, "operably linked" is intended to mean that the nucleotide sequence of interest is linked to the regulatory sequence(s) in a manner which allows for expression of the nucleotide sequence (e.g., in an *in vitro* transcription/translation system or in a host cell when the vector is introduced into the host cell). The term "regulatory sequence" is intended to include promoters, enhancers and other expression control

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elements (e.g., polyadenylation signals). Such regulatory sequences are described, for example, in Goeddel, Gene Expression Technology: Methods in Enzymology 185, Academic Press, San Diego, CA (1990). Regulatory sequences include those which direct constitutive expression of a nucleotide sequence in many types of host cell and those which direct expression of the nucleotide sequence only in certain host cells (e.g., tissue-specific regulatory sequences). It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host cell to be transformed, the level of expression of protein desired, etc.

The expression vectors of the invention can be introduced into host cells to thereby produce proteins or peptides, including fusion proteins or peptides, encoded by nucleic acids as described herein. The recombinant expression vectors of the invention can be designed for expression of a polypeptide of the invention in prokaryotic or eukaryotic cells, *e.g.*, bacterial cells such as *E. coli*, insect cells (using baculovirus expression vectors), yeast cells or mammalian cells. Suitable host cells are discussed further in Goeddel, *supra*. Alternatively, the recombinant expression vector can be transcribed and translated *in vitro*, for example using T7 promoter regulatory sequences and T7 polymerase.

Another aspect of the invention pertains to host cells into which a recombinant expression vector of the invention has been introduced. The terms "host cell" and "recombinant host cell" are used interchangeably herein. It is understood that such terms refer not only to the particular subject cell but also to the progeny or potential progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not, in fact, be identical to the parent cell, but are still included within the scope of the term as used herein. A host cell can be any prokaryotic or eukaryotic cell. For example, a nucleic acid of the invention can be expressed in bacterial cells (e.g., E. coli), insect cells, yeast or mammalian cells (such as Chinese hamster ovary cells (CHO) or COS cells). Other suitable host cells are known to those skilled in the art.

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Vector DNA can be introduced into prokaryotic or eukaryotic cells via conventional transformation or transfection techniques. As used herein, the terms "transformation" and "transfection" are intended to refer to a variety of art-recognized techniques for introducing foreign nucleic acid (e.g., DNA) into a host cell, including calcium phosphate or calcium chloride co-precipitation, DEAE-dextran-mediated transfection, lipofection, or electroporation. Suitable methods for transforming or transfecting host cells can be found in Sambrook, et al. (supra), and other laboratory manuals.

A host cell of the invention, such as a prokaryotic or eukaryotic host cell in culture, can be used to produce (*i.e.*, express) a polypeptide of the invention. Accordingly, the invention further provides methods for producing a polypeptide using the host cells of the invention. In one embodiment, the method comprises culturing the host cell of the invention (into which a recombinant expression vector encoding a polypeptide of the invention has been introduced) in a suitable medium such that the polypeptide is produced. In another embodiment, the method further comprises isolating the polypeptide from the medium or the host cell.

The host cells of the invention can also be used to produce nonhuman transgenic animals. For example, in one embodiment, a host cell of the invention is a fertilized oocyte or an embryonic stem cell into which a nucleic acid of the invention has been introduced. Such host cells can then be used to create non-human transgenic animals in which exogenous nucleotide sequences have been introduced into their genome or homologous recombinant animals in which endogenous nucleotide sequences have been altered. Such animals are useful for studying the function and/or activity of the nucleotide sequence and polypeptide encoded by the sequence and for identifying and/or evaluating modulators of their activity. As used herein, a "transgenic animal" is a non-human animal, preferably a mammal, more preferably a rodent such as a rat or mouse, in which one or more of the cells of the animal includes a transgene. Other examples of transgenic animals include non-human primates, sheep, dogs, cows, goats, chickens, amphibians, etc. A transgene is exogenous DNA which is integrated into the

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genome of a cell from which a transgenic animal develops and which remains in the genome of the mature animal, thereby directing the expression of an encoded gene product in one or more cell types or tissues of the transgenic animal. As used herein, an "homologous recombinant animal" is a non-human animal, preferably a mammal, more preferably a mouse, in which an endogenous gene has been altered by homologous recombination between the endogenous gene and an exogenous DNA molecule introduced into a cell of the animal, *e.g.*, an embryonic cell of the animal, prior to development of the animal.

A transgenic animal of the invention can be created by introducing a nucleic acid of the invention into the male pronuclei of a fertilized oocyte, e.g., by microinjection, retroviral infection, and allowing the oocyte to develop in a pseudopregnant female foster animal. The sequence can be introduced as a transgene into the genome of a non-human animal. Intronic sequences and polyadenylation signals can also be included in the transgene to increase the efficiency of expression of the transgene. A tissue-specific regulatory sequence(s) can be operably linked to the transgene to direct expression of a polypeptide in particular cells. Methods for generating transgenic animals via embryo manipulation and microinjection, particularly animals such as mice, have become conventional in the art and are described, for example, in U.S. Patent Nos. 4,736,866 and 4,870,009, U.S. Patent No. 4,873,191 and in Hogan, Manipulating the Mouse Embryo (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1986). Similar methods are used for production of other transgenic animals. A transgenic founder animal can be identified based upon the presence of the transgene in its genome and/or expression of mRNA in tissues or cells of the animals. A transgenic founder animal can then be used to breed additional animals carrying the transgene. Moreover, transgenic animals carrying a transgene encoding the transgene can further be bred to other transgenic animals carrying other transgenes.

The invention also relates to the use of the variant and reference gene products to guide efforts to identify the causative mutation for vascular diseases or to identify or synthesize agents useful in the treatment of vascular diseases, e.g., CAD and MI.

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Amino acids that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (Cunningham et al., Science, 244:1081-1085 (1989)). The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for biological activity in vitro, or in vitro activity. Sites that are critical for polypeptide activity can also be determined by structural analysis such as crystallization, nuclear magnetic resonance or photoaffinity labeling (Smith et al., J. Mol. Biol., 224:899-904 (1992); de Vos et al. Science, 255:306-312 (1992)).

Another aspect of the invention pertains to monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of proteins of the invention in clinical trials. An exemplary method for detecting the presence or absence of proteins or nucleic acids of the invention in a biological sample involves obtaining a biological sample from a test subject and contacting the biological sample with a compound or an agent capable of detecting the protein, or nucleic acid (e.g., mRNA, genomic DNA) that encodes the protein, such that the presence of the protein or nucleic acid is detected in the biological sample. A preferred agent for detecting mRNA or genomic DNA is a labeled nucleic acid probe capable of hybridizing to mRNA or genomic DNA sequences described herein, preferably in an allele-specific manner. The nucleic acid probe can be, for example, a full-length nucleic acid, or a portion thereof, such as an oligonucleotide of at least 15, 30, 50, 100, 250 or 500 nucleotides in length and sufficient to specifically hybridize under stringent conditions to appropriate mRNA or genomic DNA. Other suitable probes for use in the diagnostic assays of the invention are described herein.

The invention also encompasses kits for detecting the presence of proteins or nucleic acid molecules of the invention in a biological sample. For example, the kit can comprise a labeled compound or agent (e.g., nucleic acid probe) capable of detecting protein or mRNA in a biological sample; means for determining the amount of protein or mRNA in the sample; and means for comparing the amount of protein or mRNA in the sample with a standard. The compound or agent can be packaged in a suitable

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container. The kit can further comprise instructions for using the kit to detect protein or nucleic acid.

The following Examples are offered for the purpose of illustrating the present invention and are not to be construed to limit the scope of this invention. The teachings of all references cited herein are hereby incorporated herein by reference.

EXAMPLES

Identification of Single Nucleotide Polymorphisms

The polymorphisms shown in the Table were identified by resequencing of target sequences from individuals of diverse ethnic and geographic backgrounds by hybridization to probes immobilized to microfabricated arrays. The strategy and principles for design and use of such arrays are generally described in WO 95/11995.

A typical probe array used in this analysis has two groups of four sets of probes that respectively tile both strands of a reference sequence. A first probe set comprises a plurality of probes exhibiting perfect complementarily with one of the reference sequences. Each probe in the first probe set has an interrogation position that corresponds to a nucleotide in the reference sequence. That is, the interrogation position is aligned with the corresponding nucleotide in the reference sequence, when the probe and reference sequence are aligned to maximize complementarily between the two. For each probe in the first set, there are three corresponding probes from three additional probe sets. Thus, there are four probes corresponding to each nucleotide in the reference sequence. The probes from the three additional probe sets are identical to the corresponding probe from the first probe set except at the interrogation position, which occurs in the same position in each of the four corresponding probes from the four probe sets, and is occupied by a different nucleotide in the four probe sets. In the present analysis, probes were 25 nucleotides long. Arrays tiled for multiple different references sequences were included on the same substrate.

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Publicly available sequences for a given gene were assembled into Gap4 (http://www.biozentrum.unibas.ch/~biocomp/staden/Overview.html). PCR primers covering each exon were designed using Primer 3 (http://www-genome.wi.mit.edu/cgi-bin/primer/primer3.cgi). Primers were not designed in regions where there were sequence discrepancies between reads. Genomic DNA was amplified in at least 50 individuals using 2.5 pmol each primer, 1.5 mM MgCl₂, 100 μM dNTPs, 0.75 μM AmpliTaq GOLD polymerase, and 19 ng DNA in a 15 μ1 reaction. Reactions were assembled using a PACKARD MultiPROBE robotic pipetting station and then put in MJ 96-well tetrad thermocyclers (96°C for 10 minutes, followed by 35 cycles of 96°C for 30 seconds, 59°C for 2 minutes, and 72°C for 2 minutes). A subset of the PCR assays for each individual were run on 3% NuSieve gels in 0.5X TBE to confirm that the reaction worked.

For a given DNA, 5 μ l (about 50 ng) of each PCR or RT-PCR product were pooled (Final volume = 150-200 μ l). The products were purified using QiaQuick PCR purification from Qiagen. The samples were eluted once in 35 μ l sterile water and 4 μ l 10X One-Phor-All buffer (Pharmacia). The pooled samples were digested with 0.2 μ DNaseI (Promega) for 10 minutes at 37°C and then labeled with 0.5 nmols biotin-N6-ddATP and 15 μ Terminal Transferase (GibcoBRL Life Technology) for 60 minutes at 37°C. Both fragmentation and labeling reactions were terminated by incubating the pooled sample for 15 minutes at 100°C.

Low-density DNA chips (Affymetrix,CA) were hybridized following the manufacturer's instructions. Briefly, the hybridization cocktail consisted of 3M TMACl, 10 mM Tris pH 7.8, 0.01% Triton X-100, 100 mg/ml herring sperm DNA (Gibco BRL), 200 pM control biotin-labeled oligo. The processed PCR products were denatured for 7 minutes at 100°C and then added to prewarmed (37°C) hybridization solution. The chips were hybridized overnight at 44°C. Chips were washed in 1X SSPET and 6X SSPET followed by staining with 2 µg/ml SARPE and 0.5 mg/ml acetylated BSA in 200 µl of 6X SSPET for 8 minutes at room temperature. Chips were scanned using a Molecular Dynamics scanner.

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Chip image files were analyzed using Ulysses (Affymetrix, CA) which uses four algorithms to identify potential polymorphisms. Candidate polymorphisms were visually inspected and assigned a confidence value: high confidence candidates displayed all three genotypes, while likely candidates showed only two genotypes (homozygous for reference sequence and heterozygous for reference and variant). Some of the candidate polymorphisms were confirmed by ABI sequencing. Identified polymorphisms were compared to several databases to determine if they were novel. Results are shown in the Table.

Association of Thrombospondin Gene Polymorphisms with Vascular Disease

To determine pivotal genes associated with premature coronary artery disease, we analyzed DNA from 347 patients with MI or coronary revascularization before age 40 (men) or 45 (women) and 422 general population controls. Cases were drawn (one per family) from a retrospective collection of sibling pairs with premature CAD. Controls were ascertained through random-digit dialing. Both cases and controls were Caucasian. A complete database of phenotypic and laboratory variables for the affected patients afforded logistic regression to control for age, diabetes, body mass index, gender.

Thrombospondin (TSP) 4 and 1 emerged as important SNPs associated with premature CAD and MI. For CAD, 148 of 347 patients carried at least one copy of the TSP-4 variant compared with 142 of 422 control subjects; adjusted odds ratio 1.47, p=0.01. For premature MI, the association was even stronger: 91 of 187 cases vs. 142 of 422 controls had the variant; adjusted odds ratio 2.08, p=0.0003. The TSP-1 SNP was rare. Nonetheless, homozygosity for the variant allele gave an adjusted odds ratio of 9.5, p=.04.

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NIAF-12926 N67439 S87 DRD1, dopamine receptor D1 NIAF-12927 N67439 1279 DRD1, dopamine receptor D1 G2 NIAF-12928 N67439 1370 DRD1, dopamine receptor D1 G3 NIAF-12939 N67439 1350 DRD1, dopamine receptor D1 T3 NIAF-12933 N67439 1242 DRD1, dopamine receptor D1 T4 NIAF-12933 N67439 1242 DRD1, dopamine receptor D1 T5 NIAF-1294 N67439 1441 DRD1, dopamine receptor D1 G4 NIAF-12960 N67439 1441 DRD1, dopamine receptor D1 G5 NIAF-12961 N67439 1441 DRD1, dopamine receptor D1 G5 NIAF-12962 N67439 1465 DRD1, dopamine receptor D1 T5 NIAF-12963 N67439 264 DRD1, dopamine receptor D1 T5 NIAF-12965 N67439 264 DRD1, dopamine receptor D1 NIAF-12966 N67439 475 DRD1, dopamine receptor D1 NIAF-12966 N67439 475 DRD1, dopamine receptor D1 NIAF-12969 N67439 475 DRD1, dopamine receptor D1 NIAF-12969 N67439 1004 DRD1, dopamine receptor D1 NIAF-12971 N67439 859 DRD1, dopamine receptor D1 NIAF-12971 N67439 N67439								-		-	_
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WIAR-12928 W67439 1370 DRD1, dopamine receptor D1 B WIAR-12929 W67439 1500 DRD1, dopamine receptor D1 T WIAR-12931 W67439 1215 DRD1, dopamine receptor D1 T WIAR-12932 W67439 1215 DRD1, dopamine receptor D1 C WIAR-12934 W67439 1242 DRD1, dopamine receptor D1 C WIAR-12960 W67439 1441 DRD1, dopamine receptor D1 C WIAR-12961 W67439 1460 DRD1, dopamine receptor D1 C WIAR-12962 W67439 195 DRD1, dopamine receptor D1 C WIAR-12964 W67439 264 DRD1, dopamine receptor D1 C WIAR-12965 W67439 551 DRD1, dopamine receptor D1 C WIAR-12966 W67439 557 DRD1, dopamine receptor D1 C WIAR-12966 W67439 557 DRD1, dopamine receptor D1 C WIAR-12969 W67439 S57 DRD1, dopamine receptor D1 C WIAR-12969 W67439 S57 DRD1, dopamine receptor D1 C WIAR-12969 W67439 S57 DRD1, dopamine receptor D1 C WIAR-12969 W67439 S587 DRD1, dopamine receptor D1 C C WIAR-12969 W67439 S587 DRD1, dopamine receptor D1 C C C C C C C C C	-	IAF-12927	M67439	1279 DI	RD1,	receptor	GTGCAGCCAC[T/G]TCTGCTCCCG M T	D E	[II]	>	
WIAF-12929 M67439 1500 DRD1, dopamine receptor D1 A WIAF-12931 M67439 1338 DRD1, dopamine receptor D1 T WIAF-12932 M67439 1242 DRD1, dopamine receptor D1 A WIAF-12933 M67439 1242 DRD1, dopamine receptor D1 A WIAF-12960 M67439 1460 DRD1, dopamine receptor D1 C WIAF-12961 M67439 162 DRD1, dopamine receptor D1 C WIAF-12962 M67439 162 DRD1, dopamine receptor D1 D WIAF-12964 M67439 264 DRD1, dopamine receptor D1 D WIAF-12965 M67439 465 DRD1, dopamine receptor D1 D WIAF-12966 M67439 557 DRD1, dopamine receptor D1 D WIAF-12966 M67439 557 DRD1, dopamine receptor D1 D WIAF-12966 M67439 557 DRD1, dopamine receptor D1 D WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 D WI		IAF-12928	M67439	1370 [RD1,	dopamine receptor D1	GAAATCGCAG[C/T]TGCCTACATC M C	ا ا	4	>	
WIAF-12930 M67439 1338 DRD1, dopamine receptor D1 C WIAF-12932 M67439 1242 DRD1, dopamine receptor D1 C WIAF-12932 M67439 1242 DRD1, dopamine receptor D1 C WIAF-12934 M67439 1441 DRD1, dopamine receptor D1 G WIAF-12960 M67439 1440 DRD1, dopamine receptor D1 G WIAF-12961 M67439 399 DRD1, dopamine receptor D1 G WIAF-12963 M67439 162 DRD1, dopamine receptor D1 G WIAF-12964 M67439 264 DRD1, dopamine receptor D1 G WIAF-12966 M67439 465 DRD1, dopamine receptor D1 G WIAF-12966 M67439 557 DRD1, dopamine receptor D1 G WIAF-12966 M67439 557 DRD1, dopamine receptor D1 G WIAF-12966 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12970 M67439 859 DRD1, dopamine receptor D1 G WIAF-12970 M67439 R0104, dopamine receptor D1 G WIAF-12970 M67439 R0104 DRD1, dopamine receptor D1 G WIAF-12970 M67439 R0104, dopamine receptor D1 G WIAF-12971 M67439 R0104, dopamine receptor D1 G WIAF-12971 M67439		IAF-12929	M67439	1500 D	RD1,	receptor	ACCCTGTTGC[T/A] GAGTCTGTCT S T	E A	_4	_ K	
WIAF-12931 M67439 1215 DRD1, dopamine receptor D1 A		IAF-12930	M67439	1338 D	RD1,	dopamine receptor D1	TCTCCTACAA[C/T]CAAGACATCG S C	υ U		Z	T
WIAF-12932 M67439 1242 DRD1, dopamine receptor D1 A WIAF-12933 M67439 1441 DRD1, dopamine receptor D1 C WIAF-12934 M67439 1460 DRD1, dopamine receptor D1 G WIAF-12960 M67439 160 DRD1, dopamine receptor D1 G WIAF-12962 M67439 162 DRD1, dopamine receptor D1 G WIAF-12963 M67439 264 DRD1, dopamine receptor D1 G WIAF-12964 M67439 264 DRD1, dopamine receptor D1 G WIAF-12966 M67439 465 DRD1, dopamine receptor D1 G WIAF-12966 M67439 511 DRD1, dopamine receptor D1 G WIAF-12966 M67439 476 DRD1, dopamine receptor D1 G WIAF-12967 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 G WIAF-12970 M67439 1036 DRD1, dopamine receptor D1 G S WIAF-12971 M67439 BDD1, dopamin		IAF-12931	M67439	1215 D	RD1,		CACTCAACCC [C/A] GTCATCTATG S (υ	A D	Д	
WIAF-12933 M67439 1441 DRD1, dopamine receptor D1 G		IAF-12932	M67439	1242 D	RD1,	dopamine receptor D1	ACGCCGACTT [T/C] CAGAAGGTGT S	EH .	۲ų U	[24	
WIAF-12934 M67439 1460 DRD1, dopamine receptor D1 I WIAF-12960 M67439 399 DRD1, dopamine receptor D1 I WIAF-12961 M67439 162 DRD1, dopamine receptor D1 I WIAF-12962 M67439 264 DRD1, dopamine receptor D1 G WIAF-12964 M67439 264 DRD1, dopamine receptor D1 G WIAF-12965 M67439 511 DRD1, dopamine receptor D1 G WIAF-12966 M67439 557 DRD1, dopamine receptor D1 G WIAF-12968 M67439 476 DRD1, dopamine receptor D1 G WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G		IAF-12933	M67439	1441 D	RD1,	receptor	CGAGGAGGAG [G/A] GTCCTTTCGA M (ڻ ن	A G	വ	
WIAF-12960 M67439 399 DRD1, dopamine receptor D1 T WIAF-12961 M67439 162 DRD1, dopamine receptor D1 T WIAF-12962 M67439 264 DRD1, dopamine receptor D1 T WIAF-12963 M67439 264 DRD1, dopamine receptor D1 T WIAF-12965 M67439 511 DRD1, dopamine receptor D1 G WIAF-12966 M67439 557 DRD1, dopamine receptor D1 G WIAF-12967 M67439 476 DRD1, dopamine receptor D1 G WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 G WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 G		IAF-12934	M67439	1460 D	RD1,	receptor	GATCGCATGT [T/C] CCAGATCTAT M	E	E4 U	<u> </u>	
WIAF-12961 M67439 162 DRD1, dopamine receptor D1 T WIAF-12962 M67439 195 DRD1, dopamine receptor D1 Q WIAF-12963 M67439 264 DRD1, dopamine receptor D1 Q WIAF-12964 M67439 465 DRD1, dopamine receptor D1 Q WIAF-12965 M67439 551 DRD1, dopamine receptor D1 Q WIAF-12966 M67439 557 DRD1, dopamine receptor D1 Q WIAF-12967 M67439 476 DRD1, dopamine receptor D1 Q WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 Q WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 Q WIAF-12970 M67439 931 DRD1, dopamine receptor D1 Q		IAF-12960	M67439	399 D	RD1,		TGTCTCTGGC [C/T] GTGTCTGACC	υ	T A	A.	
WIAF-12962 M67439 195 DRD1, dopamine receptor D1 G WIAF-12963 M67439 264 DRD1, dopamine receptor D1 7 WIAF-12964 M67439 465 DRD1, dopamine receptor D1 6 WIAF-12965 M67439 511 DRD1, dopamine receptor D1 6 WIAF-12966 M67439 557 DRD1, dopamine receptor D1 7 WIAF-12968 M67439 476 DRD1, dopamine receptor D1 7 WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 7 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 7 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1 60pamine receptor D1		TAF-12961	M67439	162 D	RD1,	receptor	TGCCGCCAGG [C/G] AGCAACGGCA	υ	<u>ი</u>		_U
WIAF-12963 M67439 264 DRD1, dopamine receptor D1 7 WIAF-12964 M67439 465 DRD1, dopamine receptor D1 6 WIAF-12965 M67439 511 DRD1, dopamine receptor D1 6 WIAF-12966 M67439 557 DRD1, dopamine receptor D1 6 WIAF-12967 M67439 476 DRD1, dopamine receptor D1 6 WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 7 WIAF-12970 M67439 1036 DRD1, dopamine receptor D1 859 DRD1, dopamine receptor D1 WIAF-12970 M67439 931 DRD1, dopamine receptor D1		NIAF-12962	M67439	195	RD1,	receptor	GGCAGTTCGC [T/G] CTATACCAGC S	EH	v v	A	A
WIAF-12964 M67439 465 DRD1, dopamine receptor D1 WIAF-12965 M67439 511 DRD1, dopamine receptor D1 WIAF-12966 M67439 476 DRD1, dopamine receptor D1 WIAF-12967 M67439 476 DRD1, dopamine receptor D1 WIAF-12969 M67439 1004 DRD1, dopamine receptor D1 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		VIAF-12963	M67439	264 I	ORD1,		TGGGGCCCTC [A/G] CAGGTGGTCA S	Æ	<u>ა</u>	σ,	ß
WIAF-12965 M67439 511 DRD1, dopamine receptor D1 0 WIAF-12966 M67439 557 DRD1, dopamine receptor D1 3 WIAF-12967 M67439 476 DRD1, dopamine receptor D1 4 WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 4 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 4 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 4 WIAF-12971 M67439 831 DRD1, dopamine receptor D1 4		VIAF-12964	M67439	465 I	ORD1,	dopamine receptor D1	TGGCCGGTTA[C/T]TGGCCCTTTG	υ	E	>1	H
WIAF-12966 M67439 557 DRD1, dopamine receptor D1 WIAF-12967 M67439 476 DRD1, dopamine receptor D1 WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12965	M67439	511 [ORD1,	dopamine receptor D1	CTTCGACATC[A/T]TGTGCTCCAC M	Ą	EH	Σ	ü
WIAF-12967 M67439 476 DRD1, dopamine receptor D1 WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12966	M67439	557	ORD1,	dopamine receptor D1	ATCAGCGTGG [A/G] CCGCTACTGG M	Æ	ტ	А	_U
WIAF-12968 M67439 1004 DRD1, dopamine receptor D1 WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12967	M67439	476	DRD1,	receptor	TGGCCCTTTG [G/A] AGCGTTCTGC M	_o	A	· ·	田
WIAF-12969 M67439 1036 DRD1, dopamine receptor D1 WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12968	M67439	1004	DRD1,	dopamine receptor D1	AGCCTGCGCG [C/T] TTCCATCAAG M	υ	E	Æ	Λ
WIAF-12970 M67439 859 DRD1, dopamine receptor D1 WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12969	M67439	1036	DRD1,	receptor	GGTTCTCAAG [A/C] CCCTGTCGGT M	4	υ	H	д
WIAF-12971 M67439 931 DRD1, dopamine receptor D1		WIAF-12970	M67439	859	DRD1,	receptor	CTACATCCC [G/A] TTGCCATCAT M	ტ	Æ	>	н
	DRD5u55	WIAF-12971	M67439	931	DRD1,	dopamine receptor D1	GATTTCCTCC [C/T] TGGAGAGGGC S	Ü	E	ı,	ᄓ

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G10112	WIAF-10235	304111	1471	JUN, v-	n sarcoma virus 17	GCTGCTCAAG [C/T] TGGCGTCGCC	ა ე	E+	니	ы
G10113	WIAF-10253	704111	2010	JUN, v-	n sarcoma virus 17	TGGAGTCCCA [G/A] GAGCGGATCA	ა ზ	4	Q	Ø
2100111	WIAF-13746	D26135	993	DGKG, d. gamma (9	cerol kinase,	CCCCAGTGGT [G/A] TACCTGAAGG	დ დ	Æ	>	Þ
21001112	WTAF-13764	D26135	2313	~	diacylglycerol kinase, (90kD)	ATGTGATGAG [A/T] GAGAAACATC	M	E	ద	S
2100211	WIAF-13918	X57206	334 t	hos	ITPKB, inositol 1,4,5- trisphosphate 3-kinase B	CCCCAAGATC [A/C] GGACAAGCCT	æ	υ	a	Ωι
G1002u2	WIAF-13925	X57206	575 t	ITPKB, trisphos	ITPKB, inositol 1,4,5- trisphosphate 3-kinase B	CCAACTCAGC [T/C] TTCCTGCATA	<u>ω</u>	Ü	Æ	Æ
G1004u1	WIAF-13567	136151	1854	PIK4CA, pho kinase, cato polypeptide	phosphatidylinositol 4- catalytic, alpha ide	GCCGCTCAGA [C/T] TCCGAGGATG	ω υ	E	Д	А
G1006u1	WIAF-12375	HT2690	858	PRKCA,	protein kinase C, alpha	GGTACAAGTT [G/A] CTTAACCAAG	ري ص	A	ᄓ	니
G1008u1	WIAF-12397	HT2136	300	300 PRKCZ,	protein kinase C, zeta	CIGGCCIGCC[A/G] TGTCCGGGAG	S	_ ტ	д	д
G1008u2	WIAF-12398	HT2136	246	PRKCZ,	protein kinase C, zeta	AGTGCAGGGA [T/C] GAAGGCCTCA	E S	Ü	Д	Ω
G1008u3	WIAF-12399	HT2136	504	PRKCZ,	protein kinase C, zeta	GCTGCCACGG [C/T] CTCGTCCCGC	S C	E	<u></u>	U
G1008u4	WIAF-12403	HT2136	807	807 PRKCZ,	protein kinase C, zeta	AGAAGAATGA [C/T] CAAATTTACG	S	E⊣	Д	Д
G1008u5	WIAF-12404	HT2136	1514	1514 PRKCZ,	protein kinase C, zeta	GGATTTTCTG [A/T] CATCAAGTCC	æ	E	Ω	Δ
G1008u6	WIAF-12412	HT2136	166	166 PRKCZ,	protein kinase C, zeta	CAAGTGGGTG [G/A] ACAGCGAAGG	Œ	A	_ Ω	z
G1008117	WIAF-12418	HT2136	260	560 PRKCZ,	protein kinase C, zeta	TCCCAAGAGC [C/T] TCCAGTAGAC	Σ		_ 러	ы
G1009u1	WIAF-12396	105186	2495	PTK2, kinase	PTK2 protein tyrosine 2	TCATCAACAA [G/A] ATGAAACTGG	S	4 5	×	×
G1011u1	WIAF-11988	X07876	1250	WNT2, integra	wingless-type MMTV tion site family member 2	TCCCATGTCA[C/A]CCGGATGACC	Σ	<u>ط</u> ن	- E-	z
G1011u2	WIAF-11997	X07876	788	WNT2, integra	WNT2, wingless-type MMTV 788 integration site family member 2	GACTATGGGA [T/C] CAAATTTGCC	Σ	D E	H	E+

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21011113	WIAF-12014	X07876	1338	WNT2, wingless-type MMTV 1338 integration site family member 2	TGCACACATG [C/A] AAGGCCCCCA	U	Æ	* U	
G1011u4	WIAF-13475	X07876	856	WNT2, wingless-type MMTV 856 integration site family member 2	CCTGATGAAT [C/T] TTCACAACAA	U	H	긔	Fu
G1011u5	WIAF-13476	X07876	958	WNT2, wingless-type MMTV integration site family member 2	GACATGCTGG [C/T] TGGCCATGGC	<u>ე</u>	E	н	Li
G1011u6	WIAF-13477	X07876	789	WNT2, wingless-type MMTV 789 integration site family member 2	ACTATGGGAT [C/T] AAATTTGCCC	υ s	E	н	н
G1011u7	WIAF-13478	X07876	823	WNT2, wingless-type MMTV integration site family member 2	TGCAAAGGAA [A/G]GGAAAGGAAA	Æ	ರ	r _k	U
G1012u1	WIAF-12408	HT48910	1574	WNT2B, wingless-type MMTV 1574 integration site family, member 2B	2B ATACTTGCAA [A/G] GCCCCCAAGA	S)	.	×	×
G1016a1	WIAF-12125	Z22534	793	793 ACVR1, activin A receptor, type I	type I GGCAAGGGGA[A/G]AATGTTGCCG	8	9	БД	[x]
C1.21015	WTDE-12392	Z22534	373	373 ACVR1, activin A receptor, type I	I CTGGCCAAGC[T/C]GTGGAGTGCT	<u>ლ</u>	U	ď	A
G1018u1	WIAF-12413	X74210	1150	ADCY2, adenylate cyclase 2	CAAATTGCGA [G/T] TGGGTATTAA	Σ E	ы	>	L)
G1019u1	WIAF-12394	U83867	5475	SPTAN1, spectrin, alpha, non- 5475 erythrocytic 1 (alpha-fodrin)	eggacctaac[t/c]ggcgtgcaga	Ω E-i	ט	[-1	[-
G1019u2	WIAF-12406	U83867	1223	SPTAN1, spectrin, alpha, non- 1223 erythrocytic 1 (alpha-fodrin)	GCCCTCATCA [A/G] TGCAGATGAG	M	υ	z	ω
G1019u3	WIAF-12409	U83867	3555	SPTAN1, spectrin, alpha, non- 3555 erythrocytic 1 (alpha-fodrin)	CTGAAGGTCT [1/C]ATGGCAGAGG	⊕	U_	ᄀ	ı
G1019u4	WIAF-12415	U83867	3369	SPTAN1, spectrin, alpha, non- 3369 erythrocytic 1 (alpha-fodrin)	TCCGTGAAGC [G/A] AATGAACTAC	വ ന	- A	Æ	Æ
G1019u5	WIAF-12417	U83867	5839	SPTAN1, spectrin, alpha, non- 5839 erythrocytic 1 (alpha-fodrin)	TGAGACAGAC [T/A] TCACCGTCCA	Σ		(ž.	н
G1022u1	WIAF-12393	U45945	631	ATP1B2, ATPase, Na+/K+ transporting, beta 2 polypeptide	CATGAATGTT [A/G] CCTGTGCTGG	M	_O	H	Æ

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ATP1B2, WIAF-12400 U45945 432 transpo	ATP11 432 trans	ATPase, Na+/K+ rting, beta 2 polypeptide	GCCGCCCTGG [G/A] CGCTATTACG	Ø	ro O
ARNTL, WIAF-12401 D89722 395 nuclear	ARNT	aryl hydrocarbon receptor translocator-like	AACATTAAGA [G/C] GTGCCACCAA M	ນ ຫ	ري د
ARNTL, WIAF-12407 D89722 681 nuclear	ARNT 681 nucl	aryl hydrocarbon receptor translocator-like	CTCATAGATG [C/T] AAAAACTGGA M	EH D	A V
Home WIAF-12410 U85946 731 com	Home prol	Homo sapiens brain secretory protein hSec10p (HSEC10) mRNA, complete cds.	GATAGATTTT [C/T] AGAAGTTAAA	EH C	ω
WIAR-12402 L47647 1135 CKB,	1135 CKB	creatine kinase, brain	TCGAGATGGA [A/G] CAGCGGCTGG	A G	ы
WIAF-12405 L47647 499 CKB	499 CKB	, creatine kinase, brain	GGGAGCGCCG [A/C] GCCATCGAGA S	A C	PK.
ERCC5, excomplement complement deficiency 5 (xeroder complement HT2269 335 syndrome))	ERCC com def. 5 (: 5 com	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	GGGAICGCCA [T/C] GGGAACTCAA S	El D	二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二二
HT2269	ERC Com def 5 (Cor 1221 syr	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	CCCTCCTTCT[C/T]CAAGAACTTT M	U U	
	ERC Com def 5 (ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne			
WIAF-10431 HT2269 1783 syndrome))	1783 SYI	drome))	TCTCCAACTT [G/C] TACAAATTCT	<u>ن</u> ق	ار

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G103u4	WIAF-10432	HT2269	2077	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne	ACTGAATCTG [C/A] AGGCCAGGAT	ع ن ک	A E
G103u5	WIAF-10446	HT2269	3338	ERCCS, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne 3338 syndrome))	aatttgagct [a/t] cttgataagg	ପ ସ E	ار. ادر
G103u6	WIAE-10447	HT2269	3487	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	TCAGAATCAT[C/T]TGATGGATCT	M C T	Ω [x ₁
G103u7	WIAE-10448	HT2269	3507	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	ttcaagtgaa [C/G] atgctgaaag	ن 2 پ	D H
G103u8	WIAE-10457	HT2269	1388	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne 1388 syndrome))	CTCTTGACGA[T/G]GACGAAGATG	M F G	D M
G103u9	WIAE-10458	HT2269	1362	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	CCGGACTCTT [T/C] CAGCCATTAA	υ F4 Σ	හ අ

71031110	MT P - 1 04 59	H72269	2. 7.7.	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne	ייי אמחאמחסרט (יי, יי) מחאאממאמחרט	E-			
G103u11	WIAF-10462	HT2269	3109	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	TGGAACAGAA [C/T] GAAGACAGAT			E+	Σ
6103u12	WIAE-10463	HT2269	3138	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne 3138 syndrome))	GTTTCCTGTA [T/C] TAAAGGAACT	E E	<u>U</u>	Д	, a
G103u14	WIAE-10484	H72269	3553	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome))	agaacagctg [c/t] gaaagagcca	U Z	H	A.	Λ
G103u15	WIAF-10485	HT2269	1429	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne 1429 syndrome))	GATGTGCAGA [C/T] GGGAGGGCCA	D M	Ð	E E	Σ
G103a16	WIAP-12097	HT2269	3335	ERCC5, excision repair cross- complementing rodent repair deficiency, complementation group 5 (xeroderma pigmentosum, complementation group G (Cockayne s3335 syndrome))	AAGAATTIGA [G/T] CTACTTGATA	<u>ა</u>	EH	M	Д

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ACACTTCTGA [C/T] TGCACTCCCG	GCCACCCCAT [G/T] AACCTGGAGG	GAGTCACCAC[C/T] TTCACCTTAT	ACGTACATCA [A/C] TGCCTCGACG	TCTGTACCCA[C/T]ACTCTTGTAC	AGGCAACATG [G/C] GTGACTGGAG	Tatgtgatgc [g/a] aaaggaagag	TTCATTTTCC [G/A] AATCCTGCTG	CAAGCCTACT [C/T] AACTGCTGGA	AGABAGAGGA [A/G] GAACTCAAGG	GCACTTGAAG[C/A]AGATTGAGAT	ATGCACTTGA [A/G] GCAGATTGAG	CGCTGAGCCC[T/C]GCCAAAGACT	cctcaccaac[c/t]gctcccctct	ď
ZPK, zipper (leucine) protein 203 kinase	ZPK, zipper (leucine) protein 1806 kinase	GPR37, G protein-coupled receptor 37 (endothelin receptor type B-2825 like)	C11ORF8, chromosome 11 open 926 reading frame 8	GJA1, gap junction protein, alpha 263 1, 43kD (connexin 43)	GJA1, gap junction protein, alpha 548 1, 43kD (connexin 43)	GJA1, gap junction protein, alpha 933 1, 43kD (connexin 43)	GJA1, gap junction protein, alpha 990 1, 43kD (connexin 43)	1						
U07358 20				M65188 4	M65188 1	M65188 4	M65188 2	M65188	88	M65188	M65188	M65188	M65188	
WIAF-12411					WIAF-12438		WIAF-12440	WIAF-12441	WIAF-12442	WIAF-12465	WIAF-12466	WIAF-12486	WIAF-12487	
G1030111			G1032u1	G1033u1	G1033u2	G1033u3	G1033u4	G1033u5	G1033u6	G1033u7	G1033u8	G1033u9	G1033u10	

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210331112	WTAF-12489	M65188	GJZ 1158 1,	11, gap junction protein, alpha 43kD (connexin 43)	CTAACTCCCA [T/C] GCACAGCCTT S	D F	#	耳
G1033m13	WTAF-12490	M65188	GJ.	Al, gap junction protein, alpha 43kD (connexin 43)	TGGACATGAA [T/C] TACAGCCACT	F	기	
0.0000000000000000000000000000000000000	WTAF-12491	M65188	GJ.	A1, gap junction protein, alpha 43kD (connexin 43)	CCGCAATTAC [A/G] ACAAGCAAGC	Æ	<u>ی</u> ن	A
G1033u15	WIAF-12492	M65188	1250	GJA1, gap junction protein, alpha (1250 1, 43kD (comnexin 43)	GTGGACCAGC [G/A] ACCTTCAAGC M	ď	A R	<u> </u>
G1033u16	WIAF-12496	M65188	423	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	TATTIGIGIC [I/C] GTACCCACAC	E-	U U	ω ω
G1033u17	WIAF-12503	M65188	GJZ 880 1,	11, gap junction protein, alpha 43kD (connexin 43)	CGTTAAGGAT [C/T] GGCTTAAGGG	Ũ	H	<u>κ</u>
G1033u18	WIAF-12504	M65188	855	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	AACTCTTCTA[T/C]GTTTTCTTCA	Ен	υ	A A
G1033u19	WIAF-12505	M65188	576	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	AGITCAAGTA[C/T]GGTAITGAAG	ນ	H	A A
G1033u20	WIAF-12512	M65188	GJ 1255 1,	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	CCAGCGACCT [T/G] CAAGCAGAGC	E	ט	ري در
G1033u21	WIAF-12513	M65188	1078	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	CPACAAGCAA [G/A] CAAGTGAGCA M	_O	A	A H
2010231102	WTAF-12514	M65188	GJ.	GJA1, gap junction protein, alpha 1, 43kD (connexin 43)	CAAAACTGGG[C/G]TAATTACAGT M	U	ტ	<u>ن</u> لا
G1034u1	WIAF-12443	J03544	1201	PYGB, phosphorylase, glycogen; 1201 brain	AGACCTGTGC [A/G] TACACCAACC	4	_C	A
G1034112	WTAF-12469	J03544	771	PYGB, phosphorylase, glycogen; 771 brain	GACACCCCAG [T/C] GCCCGGCTAC	H	υ	V A
21034113	WIAF-12470	J03544	1465	PYGB, phosphorylase, glycogen;	TCCACTCGGA[G/C]ATCGTGAAAC	Ŋ	υ	E E
G1034u4	WIAF-12471	J03544	1583	PYGB, phosphorylase, glycogen; 1583 brain	GGGGCTGGCC[G/A]ATACCATCGT M	_U	Æ	Д

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G1034u5	WIAF-124/2	# # 00000	4	PYGB, phosphorylase, glycogen;	S CCTACA CA CTGTACCGGA	ა დ	0	
G1034u6	WIAF-12474	J03544	7443	PYGB, phosphorylase, glycogen;		υ υ	<u>ი</u>	
G1034u7	WIAF-12508	J03544		DPYSL2, dihydropyrimidinase-like	GCAGAGGAGC [A/G] GCAGAGGATC		Q	
G1035u1	WIAF-12484	501/60	7071		ATGACGGACC [1/C] GTGTGAAG	S H	<u>д</u>	
G1035u2	WIAF-12485	097105	7407	ppyst2 dihydropyrimidinase-like	1			
G1035u3	WIAF-12511	U97105	2062	2	CCATCACCAT [C/T] GCCAACCAGA	SC	H	н
G1036u1	WIAF-12444	D88460	311	WASL, Wiskott-Aldrich syndrome- like	ACGTGGGGTC[C/T]CTGTTGCTCA	S C	ω 01	S
G1038u1	WIAF-12445	HT2746	994	PCTK2, PCTAIRE protein kinase 2	TAGAAGAAAG [G/A] TATTGCATCG	M G A	>	ы
G1039u1	WIAF-12429	HT2747	955	55 serine/threonine kinase, PCTAIRE-3	ATCCAAGAGT [C/T] GCATGTCAGC	D D	м	J.
61039112	WIAF-12458	HT2747	808	808 serine/threonine kinase, PCTAIRE-3	CACAGAAGAG [A/T] CGTGGCCCGG	4 (ω t
G1041u1	WIAF-12459	X72886	544	544 H.sapiens TYRO3 mRNA.		ع ر ت و ت ا≊	4 2	71 2
G1041u2	WIAF-12460	X72886	693	693 H.sapiens TYRO3 mRNA.) כ		. 4
G1041u3	WIAF-12502	X72886	561	561 H.sapiens TYRO3 mRNA.	AGAGCCTGGC[C/T]GACAACCTGT	,		
G1043u1	WIAF-12448	M94055	5481	Human voltage-gated sodium channel mRNA, complete cds.	el CTCTGAGTGA [G/A] GATGACTTTG	S G	ĮΉ	ы
G1043u2	WIAF-12449	M94055	5205	Human voltage-gated sodium channel mRNA, complete cds.	el TTGAGACCTT [T/C] GGCAACAGCA	E S	[St.	[±ı
G1043u3	WIAF-12450	M94055	5224	Human voltage-gated sodium channel mRNA, complete cds.	cargarctgc[c/r]rgtrccaaar	S E	i i	ы
G1043u4	WIAF-12451	M94055	5514	Human voltage-gated sodium channel 4 mRNA, complete cds.	lel AGGITITGGGA [G/A] AAGITITGAIC	S G	Œ	댎
G1043u5	WIAF-12452	M94055	5217	Human voltage-gated sodium channel mRNA, complete cds.	GCAACAGCAT [G/C] ATCTGCCTGT	<u>ی</u>	Σ	н
G1043u6	WIAF-12453	M94055	5334	Human voltage-gated sodium channel 5334 mRNA, complete cds.	nel GCTCAGTTAA.[A/G]GGAGACTGTG	S A	×	ೱ

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G1043u7	WIAF-12454	M94055	5424	Human voltage-gated sodium 5424 mRNA, complete cds.	sodium channel	TGTACATCGC [G/C] GTCATCCTGG	ა დ	υ	Æ	A
G1043u8	WIAF-12455	M94055	5322	Human voltage-gated sodium 5322 mRNA, complete cds.	sodium channel	ATCACCCTGG [A/C] AGCTCAGTTA	ν V	<u>υ</u>	_D	ט
G1043u9	WIAF-12456	M94055	1200	Human voltage-gated sodium 1200 mRNA, complete cds.	sodium channel	ATGGCTACAC [G/A] AGCTTTGACA	<u>ი</u>	A	H	E
G1043u10	WIAF-12499	M94055	1170	Human voltage-gated sodium	sodium channel	TCTGTGTAA [G/T] GCTGGTAGAA	<u>v</u>	EH	ዾ	z
G1046a1	WIAF-13187	U50352	267	ACCN1, amiloride-sensitive cat channel 1, neuronal (degenerin)	cation cin)	TCCCAGCTGT [G/A] ACCCTCTGTA	Ω D	Æ	>	>
G1046a2	WIAF-13188	U50352	282	ACCN1, amiloride-sensitive cation 282 channel 1, neuronal (degenerin)	ion	TCTGTAACCT [C/9] AATGGCTTCC	<u>ა</u>	თ	ы	ᆈ
G1046a3	WIAF-13189	U50352	315	ACCN1, amiloride-sensitive cation 315 channel 1, neuronal (degenerin)		TCACCACCAA [C/t]GACCTGTACC	ν U	٠,٠	Z	_ z
G1046a4	WIAF-13190	U50352	386	ACCN1, amiloride-sensitive cation 386 channel 1, neuronal (degenerin)		CCCCATCTGG [C/a] TGACCCCTCC	Σ Σ		Æ	Ω
		5 C C C C C C C C C C C C C C C C C C C	717	ACCNI, amiloride-sensitive cation channel 1. neuronal (degenerin)		CCCTGCGGCA [G/A] AAGGCCAACT	ა ე	Æ	o	Ø
G1046a5	WIAF-13191	HT5174S	3214	REST, RE1-silencing factor	transcription	CAGTCAAAGC [G/A]GCTAAGGGAG	ω _O	Æ	Æ	A
G1048ul	WIAE-12641	HT5174S	3199	REST, RE1-silencing factor	transcription	caaaggaagc [c/g] ttggcagtca	S	rg _	Æ	_ 4
610404	MINE 12657	HT5174S	2125	REST, RE1-silencing factor	transcription	CTCCCATGGA [G/T] ACTGCTCAGA	<u>υ</u>	_ E-	M	А
G1048u3	WIRE-12660	HT5174S	2333	REST, RE1-silencing factor	transcription	GGAACCTGTT [A/C]AGATAGAGCT	M	<u>U</u>	×	_ a
13 15 15	WTAF-12431	HT28321	658	SCNN1G, sodium channel, 58 nonvoltage-gated 1, gamma	*	ATGACACCTC[C/T]GACTGTGCCA	S C	E .	ഗ	<u></u>
6105105	WTAF-12434	HT28321	173	SCNNIG, sodium channel, 1735 nonvoltage-gated 1, gamma		AAGCCAAGGA [G/A] TGGTGGGCCT	S	Q D	四	EI
G1051u3	WIAF-12473	HT28321	40	SCNNIG, sodium channel,		AGTCCCTGTA [T/C] GGCTTTCCAG	S	EH.	H	>1

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	L	1000 C#11	9 7 8 8	SCNNIG, sodium channel,	AGTCATTTTG [T/C] ACATAAACGA	T C	þ	121
G1051u4	MT&R-10476	HT28321	975 1		GAGGAATACA [A/G] CCCATTCCTC	M A G	z	w
Glosius	WIAF-12477	HT28321	1192	SCNNIG, sodium channel, 1192 nonvoltage-gated 1, gamma	CTGCCTACTC [G/A] CTCCAGATCT	υ υ	တ	ςς.
G1053a1	WIAF-13192	HT2201	4085	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT 4085 syndrome 3)	CGTCCTCTGA [G/A] AGCTCTGTCA	М В	M	×
G1053a2	WIAF-13193	HT2201	5607	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT syndrome 3)	ACTTTGCCGA [C/T] GCCTGTCTG	U U	Д	Ð
G1053a3	WIAF-13194	HT2201	5828	SCN5A, sodium channel, voltage- gated, type V, alpha polypeptide (long (electrocardiographic) QT spandrome 3)	GAGCCCATCA [C/T] CACCACACTC	Б Б	F	Н
G1053a4	WIAF-13202	HT2201	713	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT syndrome 3)	GCGTTCACTT [T/A] CCTTCGGGAC	E E	[± ₄]	Ы
G1053a5	WIAF-13203	HT2201	6148	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT syndrome 3)	CCACAGIGAA [G/I]AICTCGCCGA	± ±	Ð	¥
G1053a6	WIAF-13204	HT2201	6217	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT syndrome 3)	GGCCTGGCTG [G/T] CCAGGACACA	го го	t	1
G1053a7	WIAF-13205	HT2201	6324	SCN5A, sodium channel, voltage-gated, type V, alpha polypeptide (long (electrocardiographic) QT 6324 syndrome 3)	AATGGGCCTC[G/A]GCCCGCGGA	g -	1	ı

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G1054n1	WIAF-12419	HT2202	SCN4A, 2252 gated,		sodium channel, voltage- type IV, alpha polypeptide	TTGGCAAGAG [C/T] TACAAGGAGT	w	U	E	လ	ß
G1054u2	WIAF-12423	HT2202	SCN4A, 4559 gated,	i	sodium channel, voltage- type IV, alpha polypeptide	TGGTCATGTT [C/T] ATCTACTCCA	w	υ	E	ĺΞι	Įī.
G1054u3	WIAF-12424	HT2202	4856	SCN4A, gated, t	sodium channel, voltage- type IV, alpha polypeptide	TCAACATGTA [C/G] ATCGCCATCA		<u></u> υ	ŭ	×	*
G1054u4	WIAF-12425	HT2202	4777	SCN4A, gated, t	sodium channel, voltage- type IV, alpha polypeptide	GTCAAGGGTG [A/G] CTGCGGCAAC	Σ	Æ	_O	Д	ტ
G1054u5	WIAF-12426	HT2202	4863	SCN4A, gated,	sodium channel, voltage- type IV, alpha polypeptide	GTACATCGCC [A/G] TCATCCTGGA	E	Æ	_O	н	٥
G1054u6	WIAF-12427	HT2202	4566	SCN4A, 4566 gated,	sodium channel, voltage- type IV, alpha polypeptide	GITCAICIAC[I/G]CCAICIICGG	Σ	EH	<u></u>	Ø	A
G1054u7	WIAF-12428	HT2202	4923	SCN4A, gated,	sodium channel, voltage- type IV, alpha polypeptide	TGGTGAAGAT [G/T] ACTTTGAGAT	Σ			Δ	7
G1054u8	WIAF-12446	HT2202	3595	SCN4A, 3595 gated,	sodium channel, voltage- type IV, alpha polypeptide	TTCTGGCTGA [T/C] CTTCAGCATC	Σ	E	ပ	н	E
G1054u9	WIAF-12447	HT2202	4203	SCN4A, gated,	sodium channel, voltage- type IV, alpha polypeptide	GGAGACAGAC [G/A] ACCAGAGCCA	Σ	<u> </u>	Æ	Д	z
G1054u10	WIAF-12495	HT2202	4811	SCN4A, gated,	sodium channel, voltage- type IV, alpha polypeptide	TCTGCTTCTT [C/A] TGCAGCTATA	×	ט	A	<u>F4</u>	<u>-</u>
G1054u11	WIAF-12497	HT2202	5555	SCN4A, gated,	sodium channel, voltage- type IV, alpha polypeptide	CAGGCAGAC [T/G] GTGCGCCCAG	<u> </u>	H	ro l	E	H
61054012	WIAF-12498	HT2202	5480	SCN4A, 5480 gated,	sodium channel, voltage- type IV, alpha polypeptide	CAGGGGACGC [C/T] GGACCCACTA	ω l	0	E	∢	4
2105911	WIAF-12432	HT33704	112		APLP1, amyloid beta (A4) precursor-like protein 1	CGCTGCTGCT [G/A] CCACTATTGC	ω_	_ ซ	Æ	ㅂ	긔
G1059u2	WIAF-12433	HT33704	140	APLP1, precur	APLP1, amyloid beta (A4) precursor-like protein 1	TCTGCGCGCG [C/T] AGCCCGCCAT	Z	Ü	E+	O/ _	*
G1059u3	WIAF-12435	HT33704	1344	APLP1, precur	APLP1, amyloid beta (A4) 1344 precursor-like protein 1	CAGCATGTGG [C/T] CGCCGTGGAT	Σ	<u>.</u>	F	4	<u>></u>
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atgagcgaaa [g/a] gtgaatgcgt	GGTTCCTGAG [A/G] GCCAAGATGG	GTGAGGCTGT [A/G] TCGGGTCTGC	CCAAGAAATT [C/G] AAGAGGAAAT	ATCAGCCTGG [T/G] GATGCTGAGG	GCCACGGGAT [C/T] GTGGAGGTTG	CTTTGGCACC [G/A] TCATCTGCAA	GGGTGTCTGT [G/A] AGTGTGTCCA	receptor CTGCTGCTTC[1/A]GCTCTTGTTC	AAACGCTGTG [C/T]ATCATCTGGT	GTGCGCTTCT [T/C] CGCCTGCCCC	ATTTCATCAC [C/G] CTGGGCACCG	TGTGCATCAT [C/T] TGGTTCTCCT	TGAACATCAT [T/A] GACATTGTGG
APLP1, amyloid beta (A4)		()			566 CCKBR, cholecystokinin B receptor CTTTGGCACC[G/A]TCATCTGCAA	607 CCKBR, cholecystokinin B receptor GGGTGTCTGT[G/A] AGTGTGTCCA	864 CCKBR, cholecystokinin B receptor	KCNA1, potassium voltage-gated channel, shaker-related subfamily, member 1 (episodic ataxia with 684 myokymia)	KCNA1, potassium voltage-gated channel, shaker-related subfamily, member 1 (episodic ataxia with 722 myokymia)	KCNA1, potassium voltage-gated channel, shaker-related subfamily, member 1 (episodic ataxia with 804 myokymia)	KCNA1, potassium voltage-gated channel, shaker-related subfamily, member 1 (episodic ataxia with 690 myokymia)	KCNA2, potassium voltage-gated channel, shaker-related subfamily, 774 member 2
168	976	178	174	221	225	56	9	88	39	7.7	8	v	7
HT33704	HT33704	HT33704	HT1418	HT1418	HT1418	HT3538	HT3538	HT3538	HT0830	HT0830	HT0830	HT0830	HT0831
WTAE-12457	MIRE-12500	WTAR-12501	WTAR-10436	WTAF-12467	WIAF-12468	WIAF-13195	WIAF-13196	WIAF-13206	WIAF-12478	WIAF-12479	WIAF-12480	WIAF-12509	WIAF-12493
	G103044	0.0000	0100010	21080112	G1060u3	G1066a1	G1066a2	G1066a3	G1067u1	G1067u2	G1067u3	21067.114	G1068ul

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G1070a1	WIAF-13197	HT27728	522	522 member 6	CACAGTGACC[T/C]GGCTCTTTT	E C	B	ద
0.000	WTAP-13201	HT27728	1244	KCNJ6, potassium inwardly- rectifying channel, subfamily J, 1244 member 6	CCCTGGAGGA [T/C] GGGTTCTACG	S H	<u> </u>	Д
	7000 E E KTM	8077041	707	KCNJ6, potassium inwardly- rectifying channel, subfamily J, member 6	ATAAATGCCC [G/A] GAGGGAATTA	S G A		
G107083	WIAF-12422	HT48672			TTCCGGGCAA [C/T] TCAGAAGAAA	S C I		z
G1073u1	WIAF-12461	HT4556	1127	KCNJI, potassium inwardly- rectifying channel, subfamily J, 1127 member 1	CACTGTGCCA [T/C] GTGCCTTTAT	E E	<u>ک</u>	E
G1074u1	WIAF-12462	HT27804	289	KCNAB2, potassium voltage-gated channel, shaker-related subfamily, 289 beta member 2	ACCTCTTCGA [T/C] ACAGCAGAAG	S I	D U	А
G1079u1	WIAF-12463	HT27383	1130	potassium channel, inwardly 1130 rectifing (GB:D50582)	ACCTGGCCGA [T/A] GAGATCCTGT	E	A	Ed
G1079u2	WIAF-12464	HT27383	1192	potassium channel, inwardly rectifing (GB:D50582)	CGTTACTCTG [T/G] GGACTACTCC	E4	<u>۵</u>	υ υ
G1079u3	WIAF-12481	HT27383	708	potassium channel, inwardly rectifing (GB:D50582)	GCTTGGCTGC [A/G] TCTTCATGAA	Æ	D H	>
G1079u4	WIAF-12482	HT27383	977	potassium channel, inwardly 779 rectifing (GB:D50582)	CGGTGATCGC[T/C]CTGCGCCACG	S H	ر ا	<
G1079u5	WIAF-12483	HT27383	276	potassium channel, inwardly 276 rectifing (GB:D50582)	GGACCCTGCC [G/A] AGCCCAGGTA	Σ Σ	A H	
G1079u6	WIAF-12510	HT27383	489	potassium channel, inwardly 489 rectifing (GB:D50582)	GTGGCTCATC [G/A] CCTTCGCCCA	υ Σ	4	E et
G1080u1	WIAF-12536	HT4412	1095	<pre>KCNJ4, potassium inwardly- rectifying channel, subfamily J, 1099 member 4</pre>	TGGACTACTC [A/G] CGTTTTCACA	8	rg .	S

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0000	WTAB_10537	HT4412	10501	KCNJ4, potassium inwardly- rectifying channel, subfamily J,	GGCCACCGCT [T/A] TGAGCCTGTG	E E	A.	[24	×
	D C C C C C C C C C C C C C C C C C C C	100000	1 0901	potassium inwardly- ying channel, subfamily J, 2	GGCCACCGCT [A/T] TGAGCCTGTG	M	H	<u> </u>	Įżi
010010		o co chin	α 4	ium channel, inwardly ying, high conductance, subunit	CGCGGGTCAC [C/T] GAGGAGGCG	<u>ນ</u>	EH	T	E-I
G1082u1	WIAF-12662	HT28319	854	potassium channel, inwardly rectifying, high conductance, alpha subunit	CTGGTGTCGC[C/T]CATCACCATC	υ Σ	E-	<u> </u>	ы
G1082u3	WIAF-12679	HT28319	471	potassium channel, inwardly rectifying, high conductance, alpha subunit	TCTCCATCGA [G/C] ACGCAGACCA	<u>გ</u>	U	四	Д
0 to	MT&F-12198	HT0383	2028	KCNB1, potassium voltage-gated channel, Shab-related subfamily, member 1	CACTCCCCAG [C/A] AAGACTGGGG	M	4	· σ	ద
010000000000000000000000000000000000000	WIAF-13199	HT0383	2033	KCNB1, potassium voltage-gated channel, Shab-related subfamily, 2033 member 1	CCCAGCAAGA [C/G] TGGGGGGCAGC	<u>ن</u> ع		H	ω
0.10 P. 8 - 8 - 8 - 8 - 8 - 8 - 8 - 8 - 8 - 8	WT&F-13200	HT0383	2321	KCNB1, potassium voltage-gated channel, Shab-related subfamily, 2321 member 1	GAGTGTGCCA [C/A] GCTTTTGGAC	Σ	ر لا	E-I	×
480 CE	WTAF-13208	HT0383	870	KCNB1, potassium voltage-gated channel, Shab-related subfamily, 870 member 1	ACAACCCCCA [G/A] CTGGCCCACG	. σ	و و	Ø	
10 E	WIAF-12516	HT0522	1503	KCNA5, potassium voltage-gated channel, shaker-related subfamily, member 5	TCCTGGGCAA [G/A] ACCTTGCAGG	w	Ð ∆		<u>M</u>
G1088u2	WIAF-12519	HT0522	1249	KCNA5, potassium voltage-gated channel, shaker-related subfamily,	CGAGCTGCTC[G/A] TGCGCTTCTT	Σ	р А	> >	Σ

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CTCTGGGTCC [G/A] CGCGGGCCAT	GTTATCCTCA [T/C] CTCCATCATC	CAACCAGCCA [G/A] TGGAGGAGGC	CATCATCTGG [T/C] TCTCCTTCGA	GTGTATTCTG [T/a] GGATTACTCC	TTCTCTACTT [C/T] GGCTTGCGGT	GTGGTCTGCA[T/C]CTTTGGCGAC	CATGATACTT [C/G] GCTGCAGGAC	r TCGTGGTCTG[C/T]ATCTTTGGCG
KCNA5, potassium voltage-gated channel, shaker-related subfamily, member 5	KCNA5, potassium voltage-gated channel, shaker-related subfamily, 1013 member 5	KCNA6, potassium voltage-gated channel, shaker-related subfamily, 1836 member 6	KCNA3, potassium voltage-gated channel, shaker-related subfamily, 843 member 3	KCNJ8, potassium inwardly- rectifying channel, subfamily J, 1280 member 8	KCNMA1, potassium large conductance calcium-activated channel, subfamily M, alpha member	KCNMA1, potassium large conductance calcium-activated channel, subfamily M, alpha member	KCNWA1, potassium large conductance calcium-activated channel, subfamily M, alpha member	KCNWA1, potassium large conductance calcium-activated channel, subfamily M, alpha member
973	1013	1836	843	1280	765	2441	2714	200
HT0522	HT0522	HT1497	HT0222	HT27381	HT2629	HT2629	HT2629	C
WIAF-12520	WIAF-12521	WIAF-12651	WIAF-12714	WIAF-13218	WIAF-12532	WIAF-12533	WIAF-12534	5
G1088u3	G1088u4	G1090u1	G1091u1	G1094a1	G1095u1	G1095u2	G1095u3	

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G1095u5	WIAF-12539	HT2629	KC CC CJ 3048 1	CNMA1, potassium large onductance calcium-activated nannel, subfamily M, alpha member	CACTCATGAG [C/T] GCGACGTACT	හ ව	E	ω	<u>w</u>
G1095u6	WIAF-12544	HT2629	2352	KCNMA1, potassium large conductance calcium-activated channel, subfamily M, alpha member 1	GGATGTTTCA [C/T] TGGTGTGCAC	<u>က</u>	E	斑	
G1095u7	WIAF-12545	HT2629	23.92	KCNMA1, potassium large conductance calcium-activated channel, subfamily M, alpha member 1	CATCCTGACT [C/T] GAAGTGAAGC	υ υ	H	ద	*
G1095u8	WIAF-12546	HT2629	2295 H	CNWAl, potassium large onductance calcium-activated hannel, subfamily M, alpha member	CTGGCAATGA [T/C] CAGATTGACA	ω Ε	ŭ	Ω	Д
G1095u9	WIAF-12548	HT2629	29 49 7001	CNMA1, potassium large onductance calcium-activated hannel, subfamily M, alpha member	agtttttgga [c/t] caagacgatg	ა ე	E	Д	Д
G1095u10	WIAF-12549	HT2629	2865	KCNWA1, potassium large conductance calcium-activated channel, subfamily M, alpha member 1	TGCACGGGAT [G/A] TTACGTCAAC	<u>ل</u> ح	4	Σ	Н
G1096u1	WIAF-12547	126318	930	PRKM8, protein kinase mitogen- 930 activated 8 (MAP kinase)	tgctggtaat [a/t] gatgcatcta	S	H	н	н
G1098u1	WIAF-12515	119711	2650	DAG1, dystroglycan 1 (dystrophin- 2650 associated glycoprotein 1)	TCTACCTGCA [C/T] ACAGTCATTC	ა ე	H	田	斑
G110u1	WIAF-10385	HT27392	230	meiosis-specific recA homolog, 230 HsLim15	CAAAGGTATA[C/T]AGATGACAAC	N D	E	O O	*
G110u2	WIAF-10397	HT27392	1050	meiosis-specific recA homolog,	CCTGAAAATG [A/G] AGCCACCTTC	M	Ü	[32]	ღ
G110u3	WIAF-10399	HT27392	674	meiosis-specific recA homolog, 674 HsLim15	TGAACATCAG [A/G] TGGAGCTACT	X X	<u></u>	Σ	_>

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G1106u1	WIAF-12647	HT5073	5781	MAP1B, microtubule-associated protein 1B	ACTATGAGAA [G/A] ATAGAGAAA	Ŋ	A M	×
G1106u2	WIAF-12648	HT5073	5916	MAP1B, microtubule-associated protein 1B	CTGAAGAGGG [C/T] GGGTACTCAT	υ	T G	Ŋ
G1106u3	WIAF-12650	HT5073	1837	MAP1B, microtubule-associated	AGACAAGCCA [G/A] TAAAAACAGA	ტ	A V	Н
G1106u4	WIAF-12653	HT5073	2476	MAP1B, microtubule-associated 2476 protein 1B	CACCACAGGA [G/A] CIGICAIGGC	ъ	A A	E
G1106u5	WIAF-12656	HT5073	3913	MAP1B, microtubule-associated protein 1B	GCCCAATGAG [A/G] TTAAAGTCTC	A	G I	Δ
G1106u6	WIAF-12667	HT5073	559	MAP1B, microtubule-associated 559 protein 1B	GATTTTCACC [G/A] ATCAAGAGAT	ტ	A D	¤
G1106u7	WIAF-12668	HT5073	570	MAP1B, microtubule-associated protein 1B	ATCAAGAGAT[C/T]GGGGAGTTAC	۵	I L	I
G1106u8	WIAF-12669	HT5073	6175	MAP1B, microtubule-associated protein 1B	TACTTCCACA [T/C] ACTGTTACGA	H	C	出.
G1106u9	WIAF-12670	HT5073	1215	MAP1B, microtubule-associated protein 1B	TCACTCTCCA [G/C] TACCTAAACA	9	ر د	н
G1106u10	WIAF-12672	HT5073	1821	MAP1B, microtubule-associated protein 1B	AGGTAATGGT [G/A] AAAAAAGACA	_D	A >	>
G1106u11	WIAF-12673	HT5073	2727	MAP1B, microtubule-associated protein 1B	GTCCTGCCGA [G/T] TCCCCTGATG M	ъ	T E	D
G1106u12	WIAF-12674	HT5073	2739	MAP1B, microtubule-associated 2739 protein 1B	CCCCTGATGA [G/A] GGAATCACTA	ß	A E	<u>H</u>
G1106u13	WIAF-12676	HT5073	3643	MAP1B, microtubule-associated 3643 protein 1B	AGATGCCACT[G/A]ATGGCAAGGA	G	A D	Z
G1106u14	WIAF-12677	HT5073	3609	MAP1B, microtubule-associated protein 1B	CACCGCTCAA [C/T] GGATTTTCTG S	บ	Z H	Z
G1106u15	WIAF-12682	HT5073	4752	MAP1B, microtubule-associated protein 1B	TTCCAGAGCC [A/T] ACAACAGATG	Æ	Ωı E⊢	ρ,
G1110u1	WIAF-12517	HT1096	1527	1527 myelin associated glycoprotein	GCGGCCTCGT [G/C] CTCACCAGCA	υ	<u>۵</u> ان	>
G1110u2	WIAF-12518	HT1096	1678	myelin associated glycoprotein	TGTGGGCGCC [G/T] TGGTCGCCTT M	υ	> E⁺	니
G1110u3	WIAF-12522	HT1096	1271	myelin associated glycoprotein	GCCGTGTCAC [C/T] CGAGGATGAT M	U	E-1	ם
G1113u1	WIAF-12523	HT2242	353	353 myelin transcription factor 1	AATTCCGATC[G/T]GATCCTCAGG M	ย	E	저다
G1116a1	WIAF-13217	HT28451	417	<pre>myelin oligodendrocyte glycoprotein (MOG)</pre>	CAAGCTTATC [G/A] AGACCCTCTC	ß	A A	S) S)
G1116a2	WIAF-13219	HT28451	913	myelin oligodendrocyte 913 glycoprotein (MOG)	GCAGATCACT [C/G] TTGGCCTCGT M	ت ت	rg U	

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G1116a3	WIAF-13220	HT28451	922	myelin oligodendrocyte	retrigacere [g/a] retreere		A	^	<u> </u>
G1120u1	WIAF-12525	HT3695	1200	1200 neurofilament, subunit H	TAGAGATAGC [T/C] GCTTACAGAA		1	Æ	Æ
G1123u1	WIAF-12542	HT2569	2269	OMG, oligodendrocyte myelin 2269 glycoprotein	CAGCTGCAAC [T/C] CTAACTATTC	S	υ	E	E-4
G1126u1	WIAF-12526	HT28354	626	PSEN2, presenilin 2 (Alzheimer 626 disease 4)	GAGCGAAGCA [T/C] GTGATCATGC	S	Ü	出	田
G1126u2	WIAF-12527	HT28354	494	PSEN2, presenilin 2 (Alzheimer 494 disease 4)	ATGGAGAA [T/C] ACTGCCCAGT	S	บ		z
G1126u3	WIAF-12528	HT28354	434	PSEN2, presenilin 2 (Alzheimer 434 disease 4)	TAATGTCGGC[C/T]GAGAGCCCCA	S C	E	A.	Æ
G1126u4	WIAF-12543	HT28354	550	PSEN2, presenilin 2 (Alzheimer 550 disease 4)	GACCCTGACC [G/A] CTATGTCTGT	ڻ ع	₹	ద	щ
G117u1	WIAF-10391	HT27765	156	GTBP, G/T mismatch-binding 156 protein	ACTTCTCACC [A/G] GGAGATTTGG	S		Д	Δι
G117u2	WIAF-10392	HT27765	420	GTBP, G/T mismatch-binding 420 protein	AACGTGCAGA [T/C] GAAGCCTTAA	S	ບ	Ω	Д
G117u3	WIAF-10407	HT27765	939	GTBP, G/T mismatch-binding 939 protein	CCCACGTTAG [T/C] GGAGGTGGTG	ST	ט	S	S
G117u4	WIAF-10411	HT27765	1622	GTBP, G/T mismatch-binding protein	CALTGITCGA [G/A] ATTTAGGACT	MG	A	~ ~	Ж
G117u5	WIAF-10412	HT27765	2405	GTBP, G/T mismatch-binding 2405 protein	GACAGCAGGG[C/T]TATAATGTAT	М	H	Ą	Δ
G117u6	WIAF-10413	HT27765	2387	GTBP, G/T mismatch-binding	AAGAGTCAGA [A/T] CCACCCAGAC	M	H	Z	н
G125u1	WIAF-10371	HT28632	1999	ATM, ataxia telangiectasia mutated (includes complementation 1999 groups A, C and D)	CAGTAAITIT [C/I] CICAICITGI	Z Z	Ð	P	တ
G125u2	WIAF-10372	HT28632	2631	ATM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	taatgaatga [C/A] attgcagata	Σ Σ	A	D	Щ
G125u3	WIAF-10373	HT28632	3084	ATM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	CAATGGAAGA [T/G] GTTCTTGAAC	E E	ß	<u> </u>	
G125u5	WIAF-10375	HT28632	4767	ATM, ataxia telangiectasia mutated (includes complementation 4767 groups A, C and D)	CACTTATACC [C/T] CTTGTGTATG	S C	E		<u> </u>

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	WIAF-10383	HT28632	8713	ATM, ataxia telangiectasia mutated (includes complementation 8713 groups A, C and D)	ATTCTTGGAT [C/T] CAGCTATTTG	M C	H	<u> </u>	S
	WIAF-10396	HT28632	1825	ATM, ataxia telangiectasia mutated (includes complementation 1825 groups A, C and D)	GACTITGGCA [C/G] TGACCACCAG	Z Z	ტ	н	Δ
	WIAF-10398	HT28632	2924	AIM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	ACTACTGCTC [A/G]GACCAATACT	M A	ტ		K
	WIAF-10405	HT28632	8967	ATM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	TTGAAGGTGT[C/T]TTCAGAAGAT	ა ე	E	>	Λ
	WIAF-10408	HT28632	6954	ATM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	CCAAACACCT [T/C] GTAGAACTCT	E E	υ	니	ь
	WIAF-10409	HT28632	6855	ATM, ataxia telangiectasia mutated (includes complementation 6855 groups A, C and D)	TTCAGGAGCC[T/C]ATCATGGCTC	E S	υ	<u> </u>	Д
	WIAF-10410	HT28632	6801	ATM, ataxia telangiectasia mutated (includes complementation groups A, C and D)	TATATTTAA [G/T] TGGCAGAAAC	<u>ა</u>	£	저	N
	WIAF-10421	HT28632	335	ATM, ataxia telangiectasia mutated (includes complementation 335 groups A, C and D)	CATTCAGATT[C/G]CAAACAAGGA	υ Σ	ප	യ	υ
	WIAF-11607	HT28632	3966	ATM, ataxia telangiectasia mutated (includes complementation 3966 groups A, C and D)	TTCCACAICT [G/A]GTGATTAGAA	ა დ	A	ㅁ	Ь
	WIAF-13130	HT28632	8642	ATM, ataxia telangiectasia mutated (includes complementation 8642 groups A, C and D)	GAGAAATATG [A/C]AGTCTTCATG	M A	U	四	Ą
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G136u1	WIAF-10388	HT3337	535	MLH1, mutL (E. coli) homolog 1 (colon cancer, nonpolyposis type 2)	aggagaaag [c/t] tttaaaaat	Σ	H	Æ	Δ
G136u2	WIAF-10389	HT3337	MI (c	<pre>MLH1, mutL (E. coli) homolog 1 (colon cancer, nonpolyposis type 2)</pre>	TTCAAAATGA [A/G]TGGTTACATA	M. A	ტ	Z	S
G144u1	WIAF-11638	HT3625	1129	FOS, v-fos FBJ murine osteosarcoma viral oncogene homolog	CCTGTGCACT [C/T] CGGTGGTCAC	Z Z	H	<u>A</u> _	ß
G1461u1	WIAF-12562	HT0329	684	pRB-binding protein	TTGCCAAGAA [G/A] TCCAAGAACC	S G	Æ	×	×
G1466u1	WIAF-12571	HT27849	2128	2128 API2, apoptosis inhibitor 2	ATGATCCATG [G/C] GTAGAACATG	Ð	ŭ	×	U
G1468u1	WIAF-12563	HT4986	1928	apoptosis inhibitor, neuronal	CCACCAGACC [A/T] GACGAGGGC	ୟ ୟ	E		
G1468u2	WIAF-12564	HT4986	3057	3057 apoptosis inhibitor, neuronal	TTTGCAATTC [C/G] TTCAAGGGAG	Σ U	ტ	H	>
G1472u1	WIAF-12565	HT28478	242	242 BAK1, BCL2-antagonist/killer 1	GGCAGGAGTG [C/T] GGAGAGCCTG	ນ ໝ	[H	Ü	υ
G1472u2	WIAF-12572	HT28478	509	BAK1, BCL2-antagonist/killer 1	TGCAGCCCAC [G/A] GCAGAGAATG	S D	A	H	단
G1473u1	WIAF-12568	HT28606	394	CASP6, caspase 6, apoptosis- related cysteine protease	GGTGTCAACT [G/C] TTAGCCACGC	ڻ ع	Ü	>	— Н
G1473u2	WIAF-12576	HT28606	411	CASP6, caspase 6, apoptosis- related cysteine protease	ACGCAGATGC [C/T] GATTGCTTTG	s O	<u> </u>	ď	4
G1479u1	WIAF-12550	X09077	711	ATR, ataxia telangiectasia and 711 Rad3 related	ACTTTATTAA [T/C]GGTTCTTACT	E.	ບ	Σ	H
G1479u2	WIAF-12551	Y09077	4303	ATR, ataxia telangiectasia and 4303 Rad3 related	TTGCGTATGC[T/C]GATAATAGCC	S	<u>ບ</u>	A	Æ
G1479u3	WIAF-12552	X09077	1894	ATR, ataxia telangiectasia and 1894 Rad3 related	ATTCTGATGA [T/C] GGCTGTTTAA	ب ا	U s	<u> </u>	Д
G1479u4	WIAF-12553	7.090X	1855	ATR, ataxia telangiectasia and 1855 Rad3 related	ATTTATGTGG [T/A] ATGCTCTCAC	S	4	ত	יט
G1479u5	WIAF-12558	Y09077	5287	ATR, ataxia telangiectasia and 5287 Rad3 related	TCATTCATTA [T/C] CATGGTGTAG	Ω [-₁		<u> </u>	<u></u>

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G1479u6	WIAF-12559	7090Y	ATR, 5539 Rad3		ataxia telangiectasia and related	CAGCTITITA [1/C] GACTCACTGA	₽	υ	7	×
G1479u7	WIAF-12569	X09077	ATR, 1540 Rad3		ataxia telangiectasia and related	ATCCTGTTAT [1/C]GAGATGTTAG	H	ט	н	н
G1479u8	WIAF-12570	X09077	2521	ATR, Rad3 r	ataxia telangiectasia and related	ATTTAATGGA [A/G] GATCCAGACA	A	rg.	M	四
G1482u1	WIAF-12560	HT27870	3176 BLM,	BLM,	Bloom syndrome	AAAATATAAC[G/A]GAATGCAGGA	ტ	ď	L	H
G1482u2	WIAF-12561	HT27870	3605 BLM,	BLM,	Bloom syndrome	GAAATAAAGC[C/A]CAAACTGTAC	ט	A	A	Ą
G1482u3	WIAF-12573	HT27870	2677 BLM,	BLM,	Bloom syndrome	TATGTATTAC [C/T] GAAAAAGCCT M	υ 	H	д	.7
G1483u1	WIAF-12597	HT1470	1910	MYBL2, 1910 viral	v-myb avian myeloblastosis oncogene homolog-like 2	GGATGAGGAT [G/A] TGAAGCTGAT	ט	ব	>	Σ
G1483u2	WIAF-12610	HT1470	244	MYBL2, 244 viral	v-myb avian myeloblastosis oncogene homolog-like 2	ATGAGGAGGA [C/T] GAGCAGCTGA	Ü	E	Д	Д
G1483u3	WIAF-12611	HT1470	1406	MYBL2, 1406 viral	v-myb avian myeloblastosis oncogene homolog-like 2	CACTGAGAAT [A/G] GCACCAGTCT	4		α	U
G1485u1	WIAF-12581	HT1432	1941	BCR,	breakpoint cluster region	TGGAGATGAG [A/G] AAATGGGTCC	A	೮	ద	ρť
G1485u2	WIAF-12582	HT1432	3144	BCR,	breakpoint cluster region	TGACCATCAA [T/C] AAGGAAGATG	H	ŭ	z	Z
G1485u3	WIAF-12583	HT1432	3777	BCR,	breakpoint cluster region	ATAACAAGGA [T/C] GTGTCGGTGA	<u> </u>	υ	Д	Д
G1485u4	WIAF-12603	HT1432	2831	BCR,	breakpoint cluster region	CAGATCAAGA [G/A] TGACATCCAG	<u>ი</u>	4	w	×
G1485u5	WIAF-12608	HT1432	4217 BCR,	BCR,	breakpoint cluster region	ATCCCTGCCC [C/T] GGACAGCAAG	ນ _	- H	д	ц
G1486u1	WIAF-12578	HT33770	1909	BRCA2,	breast cancer 2, early	ATTGATAATG [G/A] AAGCTGGCCA	<u>ი</u>	Æ		四
G1486u2	WIAF-12579	HT33770	3623	BRCA2, onset	breast cancer 2, early	AGTTTAGAAA [A/G] CCAAGCTACA	₹	Ŋ	<u>×</u>	*
G1486u3	WIAF-12586	HT33770	1341	BRCA2, onset	breast cancer 2, early	AAATGTAGCA [A/C] ATCAGAAGCC	Æ	U U	Z	===
G1486u4	WIAF-12594	HT33770	446	BRCA2,	breast cancer 2, early	CTTATAATCA [G/A] CTGGCTTCAA	ro CO	4	α	α
G1486u5	WIAF-12598	HT33770	3013	BRCA2, 3013 onset	breast cancer 2, early	ACCATGGTTT [T/C] ATATGGAGAC	Ð	ပ	ㄹ	യ

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G1486u6	WIAF-12599	HT33770	3187	BRCA2, bre 3187 onset	breast cancer 2,	early	GAAAAATA [A/T] TGATTACATG	Z	₹.	Z F	H
G1486u7	WIAF-12604	HT33770	4971	BRCA2, onset	breast cancer 2,	early	AGCATGTGAG [A/C] CCATTGAGAT	Σ	4	٦ ت	<u>~</u>
G1486u8	WIAF-12607	HT33770	4034	BRCA2, onset	breast cancer 2,	early	ATGATICIGI [C/T] GITTCAAIGT	ß	υ	T A	
G1487u1	WIAF-12584	HT27632	2536	BRCA1, onset	breast cancer 1,	early	AGTCAGTGTG[C/G]AGCATTTGAA	Æ	<u> </u> ပ	G G	r o
G1487u2	WIAF-12587	HT27632	4697	BRCA1, onset	breast cancer 1,	early	CATCTCAAGA [G/C] GAGCTCATTA	M	ט	<u>a</u>	
G1487u3	WIAF-12595	HT27632	469	BRCA1, onset	breast cancer 1,	early	TCTCCTGAAC [A/G] TCTAAAAGAT	Ж	A (G H	ا ا
G1487u4	WIAF-12600	HT27632	3667	BRCAl, onset	breast cancer 1,	early	agcgrccaga [A/G] aggagagctr	M	Ą	n X	<u> </u>
G1487u5	WIAF-12601	HT27632	3537	BRCA1, onset	breast cancer 1,	early	TATGGGAAGT [A/G] GTCATGCATC	Σ	Æ	G S	U S
G1487u6	WIAF-12602	HT27632	4956	BRCA1, onset	breast cancer 1,	early	ATCTGCCCAG [A/G] GTCCAGCTGC	Σ	Æ	Ω Ω	ט
G1487u7	WIAF-12605	HT27632	2090	BRCA1, onset	breast cancer 1,	early	AGTACAACCA[A/G]ATGCCAGTCA	Ŋ	Æ	Q U	α o
G1487u8	WIAF-12614	HT27632	233	BRCA1, onset	breast cancer 1,	early	TCTCCACAAA [G/A] TGTGACCACA	ಬ	g	A K	X
G1492u1	WIAF-12585	HT3506	3912	cell death	3912 cell death-associated kinase	nase	TCCAGGTCCG [T/C] GGCCTGGAGA	ω	E	υ N	<u>بر</u>
G1492u2	WIAF-12593	HT3506	4352	ce11	death-associated kinase	nase	TACAACACCA [A/G] TAACGGGGT	Σ	Æ	r U	S
G1492u3	WIAF-12606	HT3506	2127	cell	death-associated ki	kinase	GCAATTTGGA [C/T] ATCTCCAACA	യ	υ	- E	A C
G1492u4	WIAF-12612	HT3506	1605	cell	death-associated kinase	nase	TGAAATTTCT [C/T] AGTGAGAACA	ω	ט	E E	- L1
G1494u1	WIAF-12589	HT28507	366	cel1	death-inducing protein Bik	ein Bik	TTCACCACAC [T/C] TAAGGAGAAC	Σ	€+.	U U	I I
G1495u1	WIAF-12580	HT27803	759	CSE1L, (yeast h	chromosome segregation homolog)-like	gation 1	ITTCTTCCCT [G/C] ATCCTGATCT	လ	υ	υ υ	- I
G1501u1	WIAF-13502	HT1949	1181	MCC, cancer	mutated in colorectal	ıtal	CAGCAATGAC [A/C] TTCCCATCGC	Σ	Æ	υ	I I
G1501u2	WIAF-13503	HT1949	1753	MCC, mutated cancers	ced in colorectal	tal	CAGCTGAGAA [C/T]GCTGCCAAGG	Ŋ	υ	E	N
G1501u3	WIAF-13504	HT1949	2344	MCC, mutated 2344 cancers	ced in colorectal	tal	TGTCCCTAGC[T/C]GAACTCAGGA	w	EH	υ	A A
G1501u4	WIAF-13521	HT1949	445	MCC, mutated	ced in colorectal	ta Ta	AGCGAACGAC [G/A] CTTCGCTATG	w	Ŋ	4	E

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G1501u5	WIAF-13522	HT1949	1504	1504 cancers	AAAGCAATGC [T/C] GAGAGGATGA	S	υ	æ	A
G1501u6	WIAF-13527	HT1949	2511	MCC, mutated in colorectal	TTCGTGAATG [A/G] TCTAAAGCGG	M	rg .	А	უ
G1502u1	WIAF-12633	HT1547	870	CCND1, cyclin D1 (PRAD1: 870 parathyroid adenomatosis 1)	AGTGTGACCC [A/G] GACTGCCTCC	ر د د	ರ	д	Д
G1503u1	WIAF-13741	U37022	1151 CDK4,	CDK4, cyclin-dependent kinase 4	CATGCCAATT [G/A] CATCGTTCAC	D E	A	υ	X
G1503u2	WIAF-13742	U37022	1410 CDK4	CDK4, cyclin-dependent kinase 4	CTGAAGCCGA [C/T] CAGTTGGGCA	ა ე	H	Д	Д
G1503u3	WIAF-13743	U37022	1328 CDK4	CDK4, cyclin-dependent kinase 4	TATGCAACAC[C/T]TGTGGACATG	D E	H	Δ,	ᄓ
G1503u4	WIAF-13780	U37022	1194	1194 CDK4, cyclin-dependent kinase 4	TTCTGGTGAC [A/G] AGTGGTGGAA	SA	ტ	H	E
G1503u5	WIAF-13781	U37022	1443 CDK4	CDK4, cyclin-dependent kinase 4	TGATTGGGCT [G/A] CCTCCAGAGG	S	ď	П	고
G1503u6	WIAF-13787	U37022	1633 CDK4	CDK4, cyclin-dependent kinase 4	CTCTTATCTA[C/T]ATAAGGATGA	υ Σ	Ð	缸	×
G1517u1	WIAF-12618	HT1132	3894	ERBB3, v-exb-b2 avian erythroblastic leukemia viral 3894 oncogene homolog 3	CAGACCTCAG [T/C] GCCTCTCTGG	S	υ	Ø	Ø
G152u1	WIAF-11608	HT3854	1673	HSPAll, heat shock 70kD protein- like 1	GTGAGTGATG[A/C]AGGTTTGAAG	Æ	Ü	ഥ	Æ
G152u2	WIAF-11629	HT3854	1683	HSPA1L, heat shock 70kD protein- like 1	AAGGTTTGAA [G/A] GGCAAGATTA	<u>ი</u>	ď	M	×
G152u3	WIAF-11609	HT3854	1478	HSPAIL, heat shock 70kD protein- like 1	GTCACAGCCA [C/T] GGACAAGAGC	υ Σ	E	H	Σ
G152u4	WIAF-11610	HT3854	1443	HSPAIL, heat shock 70kD protein- like 1	TGACGTTTGA[C/T]ATTGATGCCA	ω Ω	[-	Ω	Д
G1520u1	WIAF-12162	HT1175	2211	DNA excision repair protein ERCC2, 5' end	TGACCGTGGA[C/T]GAGGGTGTCC	ა ა	H	Ω	Д
G1520u2	WIAF-12166	HT1175	DN2 546 5'	DNA excision repair protein ERCC2, 5' end	CCCACTGCCG [A/C] TTCTATGAGG	S	ŭ	요	ĸ
G1527u1	WIAF-12168	HT0086	577	GSTM2, glutathione S-transferase 577 M2 (muscle)	TCATCTCCCG [A/C] TTTGAGGGCT	SA	υ	<u>r</u>	<u>~</u>
G1527u2	WIAF-12169	HT0086	644	GSTM2, glutathione S-transferase 644 M2 (muscle)	ACCTGTGTTC [A/T] CAAAGATGGC	M A	H	₽	യ
G1527u3	WIAF-12171	HT0086	100	GSTM2, glutathione S-transferase	ACTCAAGCTA[C/T]GAGGAAAAGA	၁ ဒ	E	→	¥
G1527u4	WIAF-12172	HT0086	41	GSTM2, glutathione S-transferase	GGGGTACTGG [A/G] ACATCCGCGG	M A	ט	Z	Д

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G1527u5	WIAF-12173	HT0086	215	8			1666 [A/ 6] C1CACASGA1	=	¢	,		
G1527u6	WIAF-12194	HT0086	238	GSTM2, M2 (musc	glutathione S-transterase :1e)		CCCAGAGCAA [T/C] GCCATCCTGC	ß	H	U	Z	z
G1528u1	WIAF-11950	HT1811	529	GSTM3, gl	glutathione S-transferase n)		GTATATTTGA [C/G] CCCAAGTGCC	Σ	Ü	U	Д	[2]
G1528112	WIAF-11951	HT1811	GST 674 M3	M3, (brai	glutathione S-transferasen)		CAACAAGCCT [G/A] TATGCTGAGC	Σ	U	Æ	>	н
G1528113	WTAF-11989	HT1811	572	GSTM3, M3 (brai	glutathione S-transferasen)		GGCTITICAIG [I/G] GCCGTITITGA	Σ	T	ש	บ	ტ
G1528u4	WIAF-13470	HT1811	240	GSTM3, M3 (brai	glutathione S-transferase		CAGAGCAATG [C/A] CATCTTGCGC	Σ	ນ	Æ	Ą	Д
G1529u1	WIAF-14146	HT2006	797 M4	IM4,	glutathione S-transferase		TGGACGCCTT [C/T] CCAAATCTGA	လ	υ	E	[건	[E4
G153u1	WIAF-12163	HT3856	1212	1212 HSPA1B,	heat shock 70kD protein	1 TGGGGC	1 TGGGCTGGA [G/A] ACGGCCGGAG	w	U	Ø	Ħ	臼
G153u2	WIAF-12182	HT3856	676	676 HSPA1B,	heat shock 70kD protein	Н	GGCCGGGGAC[A/G]CCCACCTGGG	Σ	A	ರ	₽	Æ
G153u3	WIAF-12183	HT3856	1695	1695 HSPA1B,	heat shock 70kD protein 1	1 TCAGCG	TCAGCGAGGC[C/G]GACAAGAAGA	N N	υ	Ŋ	Æ	Ø
G153u4	WIAF-12189	HT3856	330	330 HSPA1B,	heat shock 70kD protein 1 ACAAGGGGGA[G/C]ACCAAGGCAT	1 ACAAGG	GGGA [G/C]ACCAAGGCAT	Σ	ט	บ	ш	Ω
G153u5	WIAF-12190	HT3856	1053	1053 HSPA1B,	heat shock 70kD protein 1	1 AGCTGC	AGCTGCTGCA [A/G] GACTTCTTCA	ω	Æ	U	Ø	α
G1530u1	WIAF-11964	HT3010	GS' 673 M5	GSTM5, M5	glutathione S-transferase		ATTCCTCCGA [G/A] GTCTTTTGTT	Σ	υ	₹	හ	ഗ
G1530u2	WIAF-11995	HT3010	GS. 593 M5	GSTM5, M5	glutathione S-transferase		GACGCCTTCC [T/C] AAACTTGAAG	Æ	E	ပ	ы	Ωι
G1530u3	WIAF-13473	HT3010	693	GSTM5, M5	glutathione S-transferase		TTGGAAAGTC[A/G]GCTACATGGA	Ω	A	ซ	တ	တ
G1533u1	WIAF-13458	HT27460	543	GSTT2, 543 theta 2	glutathione S-transferase		CTCTCGGCTA[C/T]GAACTGTTTG	S	೮	E-I	×	<u>></u>
(21 533112	WIAF-13460	HT27460	417	GSTT2, 417 theta 2	glutathione S-transferase		GGACTGCCAT [G/A] GACCAGGCCC	Σ		∢	Σ	н
G1533u3	WIAF-13461	HT27460	359	GSTT2, 359 theta 2	glutathione S-transferase		CAGGTGTTGG [G/A] GCCACTCATT	Σ	ບ	Æ	<u></u>	ш
G1533u4	WIAF-13462	HT27460	363	GSTT2, theta 2	glutathione S-transferase		TGTTGGGGCC[A/C]CTCATTGGGG	<u>S</u>	K	೮	Д	Д
G1533u5	WIAF-13463	HT27460	385	GSTT2, 385 theta 2	glutathione S-transferase		CCAGGTGCCC [G/A] AGGAGAAGGT	Σ	ტ	ď	[22]	×
G1535u1	WIAF-11952	HT0436	517	517 HCK, he	hemopoietic cell kinase	ECGCGT	CCGCGTTGAC [T/C] CTCTGGAGAC	Σ	Е	Ü	Ω	д

G1535u2	WIAF-12013	HT0436	783 HCK,		hemopoietic cell kinase	TGGACCACTA [C/T] AAGAAGGGGA	S	E4	H	¥
G1535u3	WIAF-13464	HT0436	357	57 HCK,	hemopoietic cell kinase	TCATCGTGGT [T/C] GCCCTGTATG	S	O C	>	Þ
G1535u4	WIAF-13465	HT0436	387	387 HCK,	hemopoietic cell kinase	CCATTCACCA [C/T] GAAGACCTCA	S	E+	H	н
G1535u5	WIAF-13466	HT0436	471	471 HCK,	hemopoietic cell kinase	CCCTGGCCAC [C/G] CGGAAGGAGG	ა ე	U	터	H
G1535u6	WIAF-13467	HT0436	240	240 HCK,	hemopoietic cell kinase	CCAGCGCCAG [C/T] CCACACTGTC	w D	E-	Ω	ഗ
G1535u7	WIAF-13468	HT0436	394	94 HCK,	hemopoietic cell kinase	CCACGAAGAC [C/T] TCAGCTTCCA	Σ U	EH .	니	ĹŦ.
, in the state of	0000 t - 3 x TW	7010	MS (C	MSH2, (colon	muts (E. coli) homolog 2 cancer, nonpolyposis type	gtgaattaag[a/g]gaaataatga	Ω 4	A G	ഥ	<u> </u>
G1537u2	WIAF-12044	004045	599	MSH2, (colon 1)	<pre>mutS (E. coli) homolog 2 cancer, nonpolyposis type</pre>	GACTGTGTGA [A/T] TTCCCTGATA	M 3	A T	[2]	Д
G1537u3	WIAF-12045	U04045	MS (C 11)	MSH2, (colon 1)	mutS (E. coli) homolog 2 cancer, nonpolyposis type	AGATATGGAT [C/T]AGGTGGAAAA	Z	اط ت	Q	*
G1537u4	WIAF-12076	U04045	MS (C	MSH2, (colon 1)	<pre>mutS (E. coli) homolog 2 cancer, nonpolyposis type</pre>	GACAGITIGA [A/I] CIGACIACII	M	E H	[XI	Д
41537115	WTAF-12077	U04045	MS (C 1878 1)	MSH2, (colon 1)	muts (E. coli) homolog 2 cancer, nonpolyposis type	TCAGCTAGAT [G/A] CTGTTGTCAG	Σ.		4	H
G1543u1	WIAF-13856	000119	553	MOS, 53 viral	v-mos Moloney murine sarcoma oncogene homolog	GAGTITCIGG [G/T] CIGAGCICAA	E	E-I	4	σ.
G1543u2	WIAF-13857	J00119	621	MOS, viral	v-mos Moloney murine sarcoma oncogene homolog	GCACGCGCAC [G/A] CCCGCAGGGT	တ	<u>لا</u> ن	F	<u> </u>
G1544u1	WIAF-12018	U59464	3821	PTCH, homolog	patched (Drosophila)	CATCCCGAAT[C/T]CAGGCATCAC	Σ	C		Ω Eri
G1544u2	WIAF-12019	U59464	3618	PTCH, 3618 homolog	patched (Drosophila)	GCGIGGICCG [C/I] IICGCCAIGC	ß	E D		요 요
G1544u3	WIAF-12027	U59464	1761	PTCH, 1761 homolog	patched (Drosophila) og	ATTTTGCCAT [G/T] GTTCTGCTCA	Σ	H ق		Η Σ
G1544u4	WIAF-12029	U59464	4074	PTCH, 4074 homolog	patched (Drosophila) og	CTGCCATGGG [C/T] AGCTCCGTGC	S	U U		ro O

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G1544u5	WIAF-12043	U59464	3845	PTCH, patched (Drosophila) 3845 homolog	CCCTCGAACC[C/T]GAGACAGCAG	Ŭ ∑	Ę	д	Ц
G1544u6	WIAF-12056	U59464	1433	PTCH, patched (Drosophila) 1433 homolog	CTGCTGGTTG [C/T] ACTGTCAGTG	υ Σ	H	A	D
G1544u7	WIAF-12058	U59464	3298	PTCH, patched (Drosophila) 3298 homolog	CACCGTTCAC [G/C] TTGCTTTGGC	დ ∑	บ	>	ц
G1544u8	WIAF-12062	U59464	3986	PTCH, patched (Drosophila) 3986 homolog	TCTACTGAAG [G/A] GCATTCTGGC	ω E	Æ		田
G1544u9	WIAF-13489	U59464	1665	PTCH, patched (Drosophila)	CCATCAGCAA [T/C] GTCACAGCCT	S	C	Z	Z
G1544u10	WIAF-13490	U59464	2396	PTCH, patched (Drosophila) 2396 homolog	AAATACTTTT [C/T] TTTCTACAAC	U E	🖽	ß	Ēυ
G1544u11	WIAF-13491	U59464	2199	PTCH, patched (Drosophila)	GGACACTCTC [A/G] TCTTTTGCTG	α 4	უ	w	S
G1544u12	WIAF-13492	U59464	2222	PTCH, patched (Drosophila) 2222 homolog	AAGCACTATG [C/T] TCCTTCCTC	υ Σ	E	Æ	>
G1544u13	WIAF-13500	U59464	1686	PTCH, patched (Drosophila)	TCTTCATGGC[C/T]GCGTTAATCC	ა ე	⊟	Æ	Ą
G1545u1	WIAF-12032	HT0473	RAG1, 1835 gene	RAG1, recombination activating gene 1	GGACATGGAA [G/A] AAGACATCTT	e E	Æ	臼	×
G1545u2	WIAF-12035	HT0473	2519	RAG1, recombination activating 2519 gene 1	TGACATTGGC [A/G] ATGCAGCTGA	M	ŋ	z	А
G1545u3	WIAF-12046	HT0473	3045	RAG1, recombination activating gene 1	CGGAAAATGA [A/G] TGCCAGGCAG	4 E	ט	Z	w
G1545u4	WIAF-12047	HT0473	3146	RAG1, recombination activating 3146 gene 1	TCATAATGCA [T/C] TAAAAACCTC	S	ŭ	L	ᄓ
G1545u5	WIAF-12075	HT0473	2513	RAG1, recombination activating 2513 gene 1	CCACTGTGAC [A/T] TTGGCAATGC	Α	E	Н	ĹΣų
G1545u6	WIAF-13484	HT0473	1322	RAG1, recombination activating 1322 gene 1	GTCGCTGACT[C/T]GGAGAGCTCA	ນ ສ	[-4	PK	×
G1545u7	WIAF-13494	HT0473	2571	RAG1, recombination activating 2571 gene 1	GAAGTGTATA [A/G] GAATCCCAAT	M	ღ	×	ĸ
G1545u8	WIAF-13498	HT0473	1018	RAG1, recombination activating 1018 gene 1	TTCTGGCTGA [C/A] CCTGTGGAGA	υ Έ	A	Д	ы
G1545u9	WIAF-13499	HT0473	2782	RAG1, recombination activating 2782 gene 1	ATCTTTACCT [G/C] AAGATGAAAC	დ ე	υ	ı	ц
G1548u1	WIAF-12015	HT4999	133	IFI27, interferon, alpha- inducible protein 27	CICIGCCGIA [G/A] TITIGCCCCI	υ Σ	A.	>	н
G1548u2	WIAF-13482	HT4999	380	IFI27, interferon, alpha- 380 inducible protein 27	ATCCTGGGCT [C/T] CATTGGGTCT	υ Σ	T	Ø	ĺ±ι
G1548u3	WIAF-13483	HT4999	135	IFI27, interferon, alpha- 135 inducible protein 27	CIGCCGIAGT [T/C] TIGCCCCIGG	S	ŭ	>	Λ

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AGCTGGATGT [G/A] CCTGTGGTAA	CGGCTTCGGC[C/T]TCTCCAACTA	GCCGGGGCCA [C/T] GTGAGATTCC	TGTACGGCTT [C/T] GGCCTCTCCA	TGATGGGCAA[A/G]CAGCTGGAGA	GCAAAGTCCC[T/G]TTTAACAAAA	TGGCATACAT[C/T]GAGCGGAAGA	AAAGGCTTGG [C/T] GCTGGGCAGT	AGCCACAGAA [G/A] CCATGGGATA	CCTGCTATTT [A/T] AAAGACTTCT	acaaagttga [c/t] ttagaagaga	TCATGAGCTT [T/C] GGTATCCTTA	TCATGTAACA [A/G] AAAATCAAAT	aaaaaatcaa [a/G] tctaatagat	TTACCATGTA [A/G]AGTAAGTAAT
HC1, chromosome condensation 1	CHC1, chromosome condensation 1	HC1, chromosome condensation 1	HC1, chromosome condensation 1	HC1, chromosome condensation 1	LYN, v-yes-1 Yamaguchi sarcoma viral related oncogene homolog	LYN, v-yes-1 Yamaguchi sarcoma viral related oncogene homolog	LYN, v-yes-1 Yamaguchi sarcoma viral related oncogene homolog	LYN, v-yes-1 Yamaguchi sarcoma 996 viral related oncogene homolog	PMS1, postmeiotic segregation increased (S. cerevisiae) 1	PMS1, postmeiotic segregation increased (S. cerevisiae) 1	PMS1, postmeiotic segregation increased (S. cerevisiae) 1	PMS1, postmeiotic segregation increased (S. cerevisiae) 1	PMS1, postmeiotic segregation increased (S. cerevisiae) 1	PMS1, postmeiotic segregation 1660 increased (S. cerevisiae) 1
991 CHC1,	1271 CI	1192 CHC1,	1267 CHC1,	1657 CHC1,	LYN, 611 viral	1371 V	LYN, 1059 viral	.T 966 v	P 2355 1	2231 i	P 617 i	P 1723	1732 i	E 1
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HT3962	HT3962	HT3962	HT3962	HT3962	M16038	M16038	M16038	M16038	HT4578	HT4578	HT4578	HT4578	HT4578	HT4578
WIAF-11634	WIAF-11635	WIAF-11636	WIAF-11637	WIAF-11649	WIAF-12057	WIAF-12061	WIAF-12080	WIAF-12081	WIAF-12030	WIAF-12031	WIAF-12040	WIAF-12063	WIAF-12064	WIAF-12065
G155u1	G155u2	G155u3	G155u4	G155u5	G1550u1	G1550u2	G1550u3	G1550u4	G1552u1	G1552u2	G1552u3	G1552u4	G1552u5	G1552u6

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GAACGATACA [A/G] TAGTCAAATG	TTTAGAGGAT [G/T] CAACACTACA	TITAGACGIT [1/A] TATATAAAT	AGACGTTTTA [T/C] ATAAAATGAC	ATACCAGGAG [T/C] TTCAATTACT	TTTTCTTCT [G/T] AAAATCGATG	CTCAGAAATC[C/T]TGATGACGTC	CTGCCAGGCT [G/A] CAAGGGCCAA	CACATGCCAG [T/C] GCCAATCCCC	GGGGCTCAGC [T/C] GACGGCCCTC	GACCCCTCAG [C/T] CGTGCCTGTG	
PMS1, postmeiotic segregation 1975 increased (S. cerevisiae) 1	PMS1, postmeiotic segregation 1881 increased (S. cerevisiae) 1	PMS1, postmeiotic segregation 2454 increased (S. cerevisiae) 1	PMS1, postmeiotic segregation 7 increased (S. cerevisiae) 1	PMS1, postmeiotic segregation 2557 increased (S. cerevisiae) 1	PMS1, postmeiotic segregation	ELK3, ELK3, ETS-domain protein (SRF accessory protein 2) NOTE: 1500 Symbol and name provisional.	ELK3, ELK3, ETS-domain protein (SRF accessory protein 2) NOTE: O Symbol and name provisional.	BLK3, ELK3, ETS-domain protein (SRF accessory protein 2) NOTE:	804 PDCD1, programmed cell death 1	644 PDCD1, programmed cell death 1	
HT4578 1975	HT4578 1881	HT4578 2454	HT4578 2457	HT4578 255°	HT4578 971	HT4161 150	HT4161 1380	HT4161 1436	HT28220 80	HT28220 64	
WIAF-12066	WIAF-12067	WIAF-12068	WIAF-12069	WIAF-12082	WIAF-12083	WIAF-12028	WIAF-12059	WIAF-12060	WIAF-12024	WIAF-13488	
G1552u7	G1552u8	G1552u9	G1552u10	G1552u11	G1552u12	G1554u1	G1554u2	G1554u3	G1562u1	G1562u2	

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G1563u2	WIAF-13497	HT1187	2073	EGFR, epidermal growth factor receptor (avian erythroblastic leukemia viral (v-erb-b) oncogene 2073 homolog)	ACGGATGCAC [T/A] GGGCCAGGTC S	E	E4	E
G1566u1	WIAF-12016	HT27594	235	PDCD2, programmed cell death 2	GCGCCGCTGC [C/G] TGGCCGCCCG	υ	G G	K
G1566u2	WIAF-12033	HT27594	904	904 PDCD2, programmed cell death 2	TTGGAATTCC [A/G] GGTCATGCCT M	Æ	Q U	p:
G1566u3	WIAF-12041	HT27594	331	PDCD2, programmed cell death 2	AATCAACTAC [C/T] CAGGAAAAAC M	υ	Д. E-l	_1
G1566u4	WIAF-12071	HT27594	649	649 PDCD2, programmed cell death 2	CCTGAGGTTG [T/C] GGAAAAGGAA M	E	<u>ح</u> ن	A
G1566u5	WIAF-12072	HT27594	633	633 PDCD2, programmed cell death 2	AGAAGATGAG [A/T] TTATGCCTGA	Æ	H	Ľı,
G1567u1	WIAF-12042	M95936	293	AKT2, v-akt murine thymoma viral oncogene homolog 2	GAGAGGCCGC [G/A] ACCCAACACC M	ט	A A	Q
G1572u1	WIAF-12212	HT3998	1894	proto-oncogene c-abl, tyrosine protein kinase, alt. transcript 2	TGTTCCAGGA [A/G] TCCAGTATCT S	Ą	ඩ ස	ഥ
G1572u2	WIAF-12233	HT3998	3694	proto-oncogene c-abl, tyrosine protein kinase, alt. transcript 2	AGCTTCAGAT [C/T] TGCCCGGCGA	υ	— <u>+</u> -	н
G1572u3	WIAF-12234	HT3998	3721	proto-oncogene c-abl, tyrosine protein kinase, alt. transcript 2	GCAGTGGTCC [G/A] GCGGCCACTC S	_U	۲4 ط	Д.
G1573u1	WIAF-12021	HT0642	343	CBL, Cas-Br-M (murine) ecotropic retroviral transforming sequence	TCATGGACAA [G/C] GTGGTGCGGT	_D	O M	<u>z</u>
G1573u2	WIAF-12022	HT0642	363	CBL, Cas-Br-M (murine) ecotropic retroviral transforming sequence	TTGTGTCAGA [A/T] CCCAAAGCTG	A	Zi H	
G1573u3	WIAF-12034	HT0642	2364	CBL, Cas-Br-M (murine) ecotropic 2364 retroviral transforming sequence	AATATTCAGT[C/T]CCAGGCGCCA	υ		S Fi
G1573u4	WIAF-12049	HT0642	387	CBL, Cas-Br-M (murine) ecotropic 387 retroviral transforming sequence	CTAAAGAATA [G/A] CCCACCTTAT M	_o	o,₁ ≮	ω
G1573u5	WIAF-12050	HT0642	947	CBL, Cas-Br-M (murine) ecotropic 947 retroviral transforming sequence	AACTCATCCT [G/A] GCTACATGGC M	Ŋ	4	<u>ა</u>

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TCGAGAACCT[C/T]ATGAGTCAGG	TCTTTCCAAG [T/C] GGACTCTTTC	CTCTGGATGG [T/C] GATCCTACAA	CCGGCACTCA [C/I] ITCCAITITC	AGGGGCCCAG [C/T] TTCAGCACCA	CCCAGCGGGT [C/T] AAGAGTGACA	GAAGCCCCTG [C/T] ATGAGCAGCT	GAGAGGAAGC[C/T]GATGGGGTCT	CTGCTGGCAT [G/T] GAGTACCTGG	GATGGTCTGC[C/T]CCGGCACTTC
CBL, Cas-Br-M (murine) ecotropic 2740 retroviral transforming sequence	CBL, Cas-Br-M (murine) ecotropic 661 retroviral transforming sequence	CBL, Cas-Br-M (murine) ecotropic 2569 retroviral transforming sequence	CBL, Cas-Br-M (murine) ecotropic 2006 retroviral transforming sequence	FES, feline sarcoma (Snyder-Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v-2493 fps) oncogene homolog	FES, feline sarcoma (Snyder-Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v-189 fps) oncogene homolog	FES, feline sarcoma (Snyder-Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v-1441 fps) oncogene homolog	FES, feline sarcoma (Snyder- Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v- 2202 fps) oncogene homolog	FES, feline sarcoma (Snyder-Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v-2088 fps) oncogene homolog	FES, feline sarcoma (Snyder- Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v- 1577 fps) oncogene homolog
					V24************************************				
HT0642	HT0642	HT0642	HT0642	HT1508	HT1508	HT1508	HT1508	HT1508	HT1508
WIAF-12070	WIAF-12073	WIAF-12074	WIAF-13486	WIAF-12037	WIAF-12051	WIAF-12052	WIAF-12053	WIAF-12054	WIAF-12078
G1573u6	G1573u7	G1573u8	G1573u9	G1574u1	G1574u2	G1574u3	G1574u4	G1574u5	G1574u6

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G1574u7	WIAF-13495	HT1508	н <u>н</u> н	FES, feline sarcoma (Snyder-Theilen) viral (v-fes)/Fujinami avian sarcoma (PRCII) viral (v-fps) oncogene homolog	GTGACAAGGC [T/C]AAGGACAAGT	ω Ε	Ū	শ	ď
G1575u1	WIAF-12079	HT1052	963 b	FGR, Gardner-Rasheed feline sarcoma viral (v-fgr) oncogene	TGGGCACCGG [C/T] TGCTTCGGGG	 ວ	H	Ф	ט
G1575u2	WIAF-13487	HT1052	232 h	FGR, Gardner-Rasheed feline sarcoma viral (v-fgr) oncogene homolog	CAGAAGCTAC [G/A] GGGCAGCAGA	Σ S	4	ტ	<u>ب</u>
G1585u1	WIAF-12017	HT1675	0 966	CRK, v-crk avian sarcoma virus 996 CT10 oncogene homolog	TGGATCAACA[G/A]AATCCCGATG	ა ე	Ą	Ø	ŏ
G1585u2	WIAF-12036	HT1675	446	CRK, v-crk avian sarcoma virus	ACTACAACGT [T/C] GATAGAACCA	E	ŭ		ഗ
G1587u1	WIAF-12023	HT0590	1473 F	1473 proto-oncogene dbl	GGCCAATCCA [A/G] TTTGTGGTAC	SA	G	õ	Q
G1587u2	WIAF-12025	HT0590	2549 p	2549 proto-oncogene dbl	GTCCAGGCTT [C/T] TAATGTAGAT	C M	H	ഗ	Ē
G1587u3	WIAF-12026	HT0590	2828 p	2828 proto-oncogene dbl		Σ U	EH	လ	Fi
G1587u4	WIAF-12038	HT0590	982 g	982 proto-oncogene dbl		r E	υ	田	a
G1587u5	WIAF-12039	HT0590	2343 p	2343 proto-oncogene dbl		Σ Ω	H	Ø	н
G1587u6	WIAF-12048	HT0590	683 p	683 proto-oncogene dbl	GACACTGAAG [G/A] AGCTGTCAGT	Σ Ω	ď	ប	囝
G1587u7	WIAF-12055	HT0590	2686 p	2686 proto-oncogene dbl	TTCTCTTCAG [C/T] AGAATGATGA	C N	H	α	*
G1587u8	WIAF-13485	HT0590	2136 p	2136 proto-oncogene dbl	ACTGTGAAGG [T/A] TCTGCTCTGT	S	ď	ტ	9
G1587u9	WIAF-13496	HT0590	1566 p	1566 proto-oncogene dbl	AAAATCAGAG [C/T]AACTTAAAAA	S C	H	ß	S
G159u1	WIAF-11616	HT4209	1059 h	RAD23B, RAD23 (S. cerevisiae) 1059 homolog B	AGTACTGGGG [C/T] TCCTCAGTCT	М	T	Ą	Δ
G1590u1	WIAF-13897	HT2455	1257 C	ETS2, v-ets avian erythroblastosis virus E26 oncogene homolog 2	GCCAGTCTCT[C/G]TGCCTCAATA	S	ŭ	<u></u>	гı
G1590u2	WIAF-13913	HT2455	1107 C	ETS2, v-ets avian erythroblastosis virus E26	ATTCTGGGAC [T/G] CCCAAAGACC	S	ບ	H	E
G1590u3	WIAF-13914	HT2455	1314 c	ETS2, v-ets avian erythroblastosis virus E26 1314 oncogene homolog 2	GGAGTGACCC [A/G] GTGGAGCAAG	S	<u></u> છ	д	д
G1591u1	WIAF-13924	HT2333	1 417	HRAS, v-Ha-ras Harvey rat sarcoma 417 viral oncogene homolog	TCCAGAACCA [T/C] TTTGTGGACG	S	ŭ		н
G1595u1	WIAF-12262	HT33778	1302 t	proto-oncogene l-myc, alt. 1302 transcript 1	GCATACCTCA [G/C] TGGCTACTAA	υ Σ	ŭ	တ	H

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TD/6075	W.LAF = 1.6243	015		3A, RAI				-	-	
G160u1	WIAF-11630	HT4247	069	690 homolog A		SA	U	>	>	Т
G1602u1	WIAF-14180	HT1903	1321	1321 proto-oncogene pim-1	GTCGCCGGGG [C/A] CCAGCAAATA	Σ Σ	A	Д	H	7
G1604u1	WIAF-12319	HT2788	1182	REL, v-rel avian reticuloendotheliosis viral 1182 oncogene homolog	CCTCCCAAAG[T/C]GCTGGGATTA	S	<u>U</u>		S	
G1609u1	WIAF-12358	HT33646	348	RIPK1, receptor (TNFRSF) - interacting serine-threonine 348 kinase 1	GACGCAGGGT [C/T] TCCCATGACC	S C	[-1	Δ	Δ	
G161u1	WIAF-11654	HT4251	1522	DNA repair and recombination	TATGATCCAT [C/T] TTAACTGAGG	υ Σ	E	S	ഥ	
G1610a1	WIAF-12101	HT27727	501	501 replication protein Rpa4, 30 kDa	TGCAACTCCT [G/A] CTATTAAGAC	Σ Σ	4	4	[H	
G1610a2	WIAF-12102	HT27727	554	replication protein Rpa4, 30 kDa	TACCGTGTAA [C/T] GTGAACCAGC	ა ა	<u> </u>	Z	Z	
G1610u3	WIAF-12307	HT27727	450	replication protein Rpa4, 30 kDa	TTCTGCTGCT [G/A] ATGGAGCGAG	<u>დ</u>	A A		z	
G1610u4	WIAF-12320	HT27727	1037	1037 replication protein Rpa4, 30 kDa	TGATTCATGA [G/C] TGTCCTCATC	∑ ∑	Ö	[2]	Д	
G1610u5	WIAF-12321	HT27727	857	57 replication protein Rpa4, 30 kDa	TAGAGGACAT [G/A] AACGAGTTCA	স ড	4	Σ	н	T
G1610u6	WIAF-12343	HT27727	539	539 replication protein Rpa4, 30 kDa	GAATTCAGGA [C/T] GTTGTACCGT	ຮ	H	Α		
G1630u1	WIAF-12302	HT3563	4312	DCC, deleted in colorectal carcinoma	ACTCATGAAG[C/T]AGCTTAATGC	N		δ	*	
G1632u1	WIAF-13572	HT27355	742	tumor suppressor, PDGF receptor 742 beta-like	TTTATGACAT [G/C] AAGCGGGGCT	Σ		Σ	H	
G1632u2	WIAF-13584	HT27355	1102	tumor suppressor, PDGF receptor	TGGAAGACTT[C/T]GAGACGATTG	8	ط ن	[E ₄	[II4	
G1632u3	WIAF-13601	HT27355	258	tumor suppressor, PDGF receptor 258 beta-like	AAGACGCAGT [C/T] TATCATGATG	Σ	E C	S	(II.)	
G1633u1	WIAF-13957	HT1778	1263	FER, fer (fps/fes related) tyrosine kinase (phosphoprotein NCP94)	ttcaggcara [t/c] gagatcatgt	တ	υ -	Z	z	
G1633u2	WIAF-13958	HT1778	2407	FER, fer (fps/fes related) tyrosine kinase (phosphoprotein 2407 NCP94)	TATGTTGTAT [C/T] TCGAGAGTAA	Σ	U U		Įtų	
G1634u1	WIAF-13505	HT3216	1569	BLK1, BLK1, member of BTS 1569 oncogene family	TCTCGACCCC [C/T] GTGGTGCTCT	ω O	ا ت		Ωı	

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G1634u2	WIAF-13858	HT3216	456	ELK1, BLK1, member of ETS 456 oncogene family	GGCTGTGGGG [A/G] CTACGCAAGA	Ω 4	_D	უ	ტ
G1634u3	WIAF-13859	HT3216	745	ELK1, BLK1, member of BTS 745 oncogene family	AGGCCCAGGC [G/A] GTTTGGCACG	ĭ B	ď.	ტ	S
G1638u1	WIAF-14172	HT1224	86	98 uracil-DNA glycosylase	GCTGGGACCT [G/C] TTCCACAAAT	- G	Ŋ		_
G1643u1	WIAF-13517	HT3751	629	DXS648E, DNA segment on chromosome X (unique) 648 629 expressed sequence	TACATCCCCA [G/A] TCGTGGCCCT	<u>დ</u>	Æ	Ω	Z
G1645u1	WIAF-14087	D21089	363	XPC, xeroderma pigmentosum, 363 complementation group C	AAAACCICAA [G/A] GITATAAAGG	<u>හ</u>	_		×
G1645u2	WIAF-14088	D21089	2166	XPC, xeroderma pigmentosum, 2166 complementation group C	TGCATTCCAG [G/A]GACACGTGGC	ა ე	4	ρ;	<u></u> K
G1645u3	WIAF-14089	D21089	1580	XPC, xeroderma pigmentosum, 1580 complementation group C	GGGAGCCATC[G/A]TAAGGACCCA	ت ع	Ą		斑
G1645u4	WIAF-14090	D21089	1601	XPC, xeroderma pigmentosum, 1601 complementation group C	AGCTTGCCAG [T/C] GGCATCCTCA	E	ರ	Þ	Æ
G1645u5	WIAF-14091	D21089	2920	XPC, xeroderma pigmentosum, complementation group C	CCCATTTGAG [A/C] AGCTGTGAGC	M	ت ت	<u>×</u>	Q
G1645u6	WIAF-14103	D21089	405	XPC, xeroderma pigmentosum,	ATGACCTCAG [G/A] GACTTTCCAA	ත ආ	4;	ద	ద
G1645u7	WIAF-14104	D21089	151	XPC, xeroderma pigmentosum, 151 complementation group C	GGGACGCGAA [C/G] TGCGCAGCCA	M	D.	니	Δ
G1645u8	WIAF-14105	D21089	2133	XPC, xeroderma pigmentosum, 2133 complementation group C	AAGCGGTCTA[C/T]TCCAGGGATT	ນ	H	- A	<u> </u>
G167u1	WIAF-11632	HT4579	83	PMS2L8, postmeiotic segregation increased 2-like 8	CCTATTGATC [G/A] GAAGTCAGTC	ڻ ح	Æ		α
G167u2	WIAF-11633	HT4579	219	PMS2L8, postmeiotic segregation increased 2-like 8	GAGTGGATCT[T/C]ATTGAAGTTT	S L	ŭ	니	ы
G167u3	WIAF-11644	HT4579	768	PMS2L8, postmeiotic segregation 768 increased 2-like 8	TGCCCCCTAG [T/C] GACTCCGTGT	S)	U	<u> </u>	ς,

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G167u4	WIAF-11622	HT4579	1645	PMS2L8, postmeiotic segregation 1645 increased 2-like 8	GAAAGCGCCT [G/A] AAACTGACGA		Ą	四	×
G167u5	WIAF-11645	HT4579	1512	PMS2L8, postmeiotic segregation increased 2-like 8	ACTCGGGGCA [C/T] GGCAGCACTT	ა ა	E	田	
G167u6	WIAF-11646	HT4579	1619	PMS2L8, postmeiotic segregation	TCGCAGGAAC [A/G] TGTGGACTCT	Σ V			ద
G167u7	WIAF-11647	HT4579	1432	PMS2L8, postmeiotic segregation	CGTCCTGAGA [C/T] CTCAGAAAGA	υ Σ	H		တ
G167u8	WIAF-11625	HT4579	2490	PMS2L8, postmeiotic segregation 2490 increased 2-like 8	GGACTGCTCT [T/C] AACACAAGCG	ري ا	<u></u>	— Н	- н
G167u9	WIAF-11619	HT4579	804	PMS2L8, postmeiotic segregation 804 increased 2-like 8	TGAGCTGTTC[G/C]GATGCTCTGC	න න	ָט	လ	<u> </u>
G167u10	WIAF-11623	HT4579	1555	PMS2L8, postmeiotic segregation	CATCCCAGAC [A/G] CGGGCAGTCA	E E	ro	E	4
G167u11	WIAF-11624	HT4579	2364	PMS2L8, postmeiotic segregation increased 2-like 8	CCTTCGGACC[C/T]CAGGACGTCG	ى ن	H	Ωı	д
G167u12	WIAF-11626	HT4579	2348	PMS2L8, postmeiotic segregation 2348 increased 2-like 8	ACTAGTAAAA [A/G] CTGGACCTTC	M A	ָט	Z	Ŋ
G181u1	WIAF-11697	HT48793	311	ERCC4, excision repair cross- complementing rodent repair deficiency, complementation group 4	ATATTTGCGA [C/T] AAGTAGGATA	<u>υ</u> Σ	H	Ė	Н
G181u2	WIRF-11698	HT48793	EE CC CC 295 4 295 4	ERCC4, excision repair cross-complementing rodent repair deficiency, complementation group 4	CACACAAGGT [G/C] GTGTTATATT	M	G C	ტ	ద
G181u3	WIAF-11699	HT48793	234	ERCC4, excision repair cross- complementing rodent repair deficiency, complementation group	TTGAACACCT [C/T] CCTCGCCGTG	ω O	U U		

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				ERCC4, excision repair cross- complementing rodent repair					
G181u4	WIAF-11704	HT48793	808	deficiency, complementation group	TTTGTGGCAC [C/T] AGCTTGGAGC	U N	E	ď	*
				ERCC4, excision repair cross-complementing rodent repair deficiency, complementation group					
G181u5	WIAF-11705	HT48793	640 4	4	TTCTATGACA[C/T]CTACCATGCT	ນ ≅	ㅂ	24	20
				ERCC4, excision repair cross-complementing rodent repair deficiency, complementation group	()))))) () () () () () () (<u>ح</u>	E	Д	ď
G181u6	WIAF-11670	HT48793	1117		AGAAAGCAAC [C/ 1] CAAAG1GGGA		4	4	,
G185u1	WIAF-11668	HT5122	ACV 319 IIB	ACVR2B, activin A receptor, type IIB	TCTGCAACGA [G/A] CGCTTCACTC	ω D	A	闰	ſΞI
0,50	WISE-11707	HT5122	702	ACVR2B, activin A receptor, type IIB	AGACACGGGA [G/C] TGCATCTACT	M G	บ	函	Д
ST COTE				ACVR2B, activin A receptor, type	C C C 4 4 C C C 2 C C 4 C C / E C 2 C C C C C C C C C C C C C C C C C	E	ľ	>	þ
G185u3	WIAF-11672	HT5122	812		CCTCACGGAT [T/C] ACCTCAAGGG	T	ار	4	4
G185u4	WIAF-13542	X77533	1109	ACVR2B, activin A receptor, type IIB	GGCTCCTGAG [G/A] TGCTCGAGGG	∑ ∑	Æ	>	×
0.00	12 T T T T T T T T T T T T T T T T T T T	X77533	664	ACVR2B, activin A receptor, type	TGCTGAAGAG [C/T] GACCTCACAG	ာ ၁	E-1	Ω	ß
G18711	WTAF-11669	HT97400	183	183 androgen	CCAGAGACAG [C/T] GCGACCCGGA	υ Σ	H	ద	Ü
		6 C C C C C C C C C C C C C C C C C C C	7.7	CXCR4, chemokine (C-X-C motif),	ACCTGGCCAT [C/T] GTCCACGCCA	<u>ი</u>	EH	н	н
G191ul	WIAF-101/6	AF 025375	# #	TOTATO OTO		-	ļ		
G193u1	WIAF-10178	D29984	231	cckz, recepto	AGTGCTTGAC [T/A] GACATTTACC	S	A	E	E
G193u2	WIAF-10179	D29984	190	CCR2, chemokine (C-C motif) 190 receptor 2	CATGCTGGTC [G/A] TCCTCATCTT	Σ.	_ <	>	н
G194u1	WIAF-10211	D43767	121	SCYA17, small inducible cytokine subfamily A (Cys-Cys), member 17	ACATCCACGC [A/C] GCTCGAGGGA	ري ح	Ü	A	Ą
G197n1	WIAF-10167	D50403	1515	NRAMP1, natural resistance- associated macrophage protein 1 (might include Leishmaniasis)	GGTGCTAGTC[T/C]GCGCCATCAA	E E	U	Ü	ద
G197u2	WIAF-10173	D50403	1629	NRAWD1, natural resistance- associated macrophage protein 1 (might include Leishmaniasis)	CACCTACCTG [G/C] TCTGGACCTG	<u>ა</u>	<u> </u>	>	<u>г</u>

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				ACVRIB, activin A receptor, type		_			
G20u1	WIAF-10249	U14722	896 IB		CGGTACACAG [T/C] GACAATTGAG	Ε Σ	ט	>	A
G20u2	WIAF-10250	U14722	ACT 866 IB	ACVRIB, activin A receptor, type IB	GAGCACGGGT [C/T] CCTGTTTGAT	υ Σ	₽	ß	ĹŦ4
G20113	WIAF-10251	014722	1391	ACVR1B, activin A receptor, type IB	CAGAGTTATG [A/T] GGCACTGCGG	M M	. E-	闰	Þ
G2014	WIAF-10252	U14722	AC' 1236 IB	ACVRIB, activin A receptor, type IB	TATATTGGGA [G/C]ATTGCTCGAA	υ Σ	_ ပ	印	Д
G20115	WIAF-10261	U14722	AC 518 IB	ACVR1B, activin A receptor, type IB	GAGATGTGTC [T/C] CTCCAAAGAC	E E	บ	ᄓ	Ъ
G207a1	WIAF-10516	L25259	998	Human CTLA4 counter-receptor (B7-866 2) mRNA, complete cds.	AGCTGTACTT [C/T] CAACAGTTAT	ນ ຮ	E	Д	S
G208u1	WIAF-10204	131581	85	CCR7, chemokine (C-C motif) 85 receptor 7	GGGGAAACCA [A/G] TGAAAAGCGT	M A	_D	Σ	>
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6211U1	WIAF-10191	M27533	452	i ë i	TGAAAGAAGT [G/A] GCAACGCTGT	დ ტ	A	>	Δ
131100	WTAF-11659	M28393	822		GCATCTCTGC[C/T]GAAGCCAAGG	S	E	⋖	Æ
421511	WIAF-11723	M28393	159	PRF1, perforin 1 (preforming 159 protein)	TGACCAGCCT [C/T] CGCCGCTCGG	ນ ບ	E	н	1
G215u3	WIAF-11724	M28393	96	PRF1, perforin 1 (preforming 96 protein)	CAGAGTGCAA [G/A] CGCAGCCACA	ω Θ	Æ	×	×
G215u4	WIAF-11725	M28393	1377	PRF1, perforin 1 (preforming 1377 protein)	ATAACAACCC [C/T] ATCTGGTCAG	S	EH	Д	Д
G215u5	WIAF-11726	M28393	1326	PRF1, perforin 1 (preforming 1326 protein)	TGAAGCTCTT [C/T] TTTGGTGGCC	S C	H	[E4	[14
G215u6	WIAF-11727	M28393	1076	PRF1, perforin 1 (preforming 1076 protein)	CGGCGGGAGG [C/T] ACTGAGGAGG	Σ U	H	A	>
G217u1	WIAF-11691	M31932	649	FCGR2B, Fc fragment of IgG, low affinity IIb, receptor for (CD32)	GCAGCTCTTC [A/G] CCAATGGGGA	⊗ 4	ტ	ω.	ω
G2 <u>1</u> 7u2	WIAF-11692	M31932	625	FCGR2B, Fc fragment of IgG, low 625 affinity IIb, receptor for (CD32)	TCACTGTCCA[A/G]GTGCCCAGCA	S A	U	O)	0

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G217u3	WIAF-11712	M31932	332 af	FCGR2B, Fc fragment of IgG, low affinity Ilb, receptor for (CD32)	GACTGGCCAG [A/C] CCAGCCTCAG	M	Ü	H	д
G217u4	WIAF-11713	M31932	FC 101 af	FCGR2B, Fc fragment of 1gG, low affinity IIb, receptor for (CD32)	GGCTTCTGCA [G/T] ACAGTCAAGC	ڻ Σ	H		>
G218u1	WIAF-10184	M36712		CD8B1, CD8 antigen, beta polypeptide 1 (p37)	ttttacaaat [a/g] agcagagat	N	Ü	*	*
G218u2	WIAF-10188	M36712	326 pc	CD8B1, CD8 antigen, beta 326 polypeptide 1 (p37)	GCTGTGTTTC [G/C] GGATGCAAGC	∑ છ	_ ပ	pz	Д
G218u3	WIAF-10189	M36712	196 pc	CD8B1, CD8 antigen, beta polypeptide 1 (p37)	CAGTAACATG [C/T] GCATCTACTG	υ Σ	E	ద	ט
G218u4	WIAF-10190	M36712	225 pc	CD8B1, CD8 antigen, beta 5 polypeptide 1 (p37)	AGCGCCAGGC [A/C] CCGAGCAGTG	8	Ü		₫.
G218u5	WIAF-10194	M36712	583 pc	CD8B1, CD8 antigen, beta polypeptide 1 (p37)	GGTGGCTGGC [G/A] TCCTGGTTCT	υ Σ	A	>	н
G218u6	WIAF-10208	M36712	372 pc	CD8B1, CD8 antigen, beta 372 polypeptide 1 (p37)	TGAAGCCGGA [A/G] GACAGTGGCA	S A	Ü	[2]	阳
G218u7	WIAF-10209	M36712	400 pc	CD8B1, CD8 antigen, beta	CTGCATGATC [G/T] TCGGGAGCCC	<u>ტ</u> გ	Ð	>	ſ±ι
G218u8	WIAF-10210	M36712	270 pc	CD8B1, CD8 antigen, beta 270 polypeptide 1 (p37)	TCTGGGATTC [C/T] GCAAAAGGGA	S	EH	တ	ഗ
G218a9	WIAF-10518	M36712	CI 618 pc	CD8B1, CD8 antigen, beta 618 polypeptide 1 (p37)	GAGTGGCCAT [C/G] CACCTGTGCT	Σ	හ	н	Σ
G218a10	WIAF-13223	M36712	CJ CJ 556 pv	CD8B1, CD8 antigen, beta 556 polypeptide 1 (p37)	Trefagence [a/6] Teaccettes	Æ	ڻ ن	н	>
G218a11	WIAF-13224	M36712	836 p	CD8B1, CD8 antigen, beta polypeptide 1 (p37)	CTGTGTGTGA [T/C] GTGCATGGGA	1	U		
G22u1	WIAF-10301	U86136	H 6719 p	Human telomerase-associated 6719 protein TP-1 mRNA, complete cds.	GCTGGTAACC [G/A] TCGGGCTAGA	Σ O	4	D D	н
G22u2	WIRF-10302	U86136	H 7537 p	Human telomerase-associated protein TP-1 mRNA, complete cds.	CTGATGGGAT [C/G] CTATGGAACC	Σ	<u>စ</u>	н	Σ
G22u3	WIAF-10311	U86136	H 1798	Human telomerase-associated 1798 protein TP-1 mRNA, complete cds.	ATGATGCCAT [T/C] GATGCCCTCG	ω z	D E	H	н
G22u4	WIAF-10312	U86136	2397 p	Human telomerase-associated protein TP-1 mRNA, complete cds.	CTGTCTCGG [C/T] TGGCCAAAGG	Æ	El C	₹	>

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G22u5	WIAF-10313	U86136	3289	Human telomerase-associated 3289 protein TP-1 mRNA, complete cds.	AGAAAGGGAT [A/C] ACCTGCCGCA	α, S	υ	н	
G22u6	WIAF-10314	U86136	3242 E	Human telomerase-associated 3242 protein TP-1 mRNA, complete cds.	AGAGGCCGCA [T/C] GTCGGATCTC	Ε	ŭ	υ R	1
G22u7	WIAF-10315	U86136	4482 E	Human telomerase-associated 4482 protein TP-1 mRNA, complete cds.	CCGTTTGCCT [G/A] CCTCGTCCAG	<u>ප</u>	Æ	<u>۲</u>	
G22u8	WIAF-10316	U86136	1 4363	Human telomerase-associated 4363 protein TP-1 mRNA, complete cds.	GTTTGACTGT [G/A] GACCAGCTGC	<u>ა</u>	Æ	Δ Δ	
G22u9	WIAF-10317	U86136	4230	Human telomerase-associated 4230 protein TP-1 mRNA, complete cds.	GTGTCTGAGA [G/A] ACTCCGGACC	<u>დ</u>	A	2	M
G22u10	WIAF-10318	U86136	4419	Human telomerase-associated 4419 protein TP-1 mRNA, complete cds.	GGGACTRAGA [G/C] CTGGGAAGAA	ڻ ح	υ	ω	E
G22u11	WIAF-10319	U86136	5269	Human telomerase-associated 5269 protein TP-1 mRNA, complete cds.	TCTCCGATGA [T/C] ACACTCTTTC	ω H	Ü	A	Q
G22u12	WIAF-10320	U86136	5015	Human telomerase-associated 5015 protein TP-1 mRNA, complete cds.	GCTGCTCTCC [C/T] GGAGATGGCA	υ Σ	E	ıχ.	×
G22u13	WIAF-10321	186136	5133	Human telomerase-associated 5133 protein TP-1 mRNA, complete cds.	GTGGCCTTCT[C/T]CACCAATGGG	Σ.	H	Ŋ	[ī4
G22u14	WIAF-10322	U86136	7764	Human telomerase-associated 7764 protein TP-1 mRNA, complete cds.	ACAGCCCTCC [A/G] TGTGCTACCT	Σ α	ъ	Ħ	K
G22u15	WIAF-10323	U86136	7884	Human telomerase-associated 7884 protein TP-1 mRNA, complete cds.	TGCCTGGAAC [C/T] TTGGCTGGGC	υ Σ	E .	д	L
G22u16	WIAF-10324	U86136	7744	Human telomerase-associated protein TP-1 mRNA, complete cds.	AGATTCACTC [G/A] GGCTCTGTCA	ν υ	K	S	ω.
G22u17	WIAF-10337	U86136	1018	Human telomerase-associated 1018 protein TP-1 mRNA, complete cds.	CCATTGCTGC [T/C] TTCTTGCCGG	Ω E-l	Ü	Æ	A
G22u18	WIAF-10338	U86136	1000	Human telomerase-associated 1000 protein TP-1 mRNA, complete cds.	tggccaataa [c/a] atcttggcca	Σ O	4	z	×

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G22u19	WIAF-10339	U86136	1182	Human telomerase-associated	ATGACGGACA [A/G]ATTTGCCCAG	Æ	_U	ᄶ	ex
G22u20	WIAF-10340	U86136	1939	Human telomerase-associated 1939 protein TP-1 mRNA, complete cds.	AGCAGCTTCG [T/G] ATGGCAATGA	S H	ტ	ద	ద
G22u21	WIAF-10341	U86136	1 2227	Human telomerase-associated 2227 protein TP-1 mRNA, complete cds.	TCACGAGGGC [G/A] GAGCAGGTGG	დ დ	Æ	Æ	4
G22u22	WIAF-10342	U86136	2776	Human telomerase-associated 2776 protein TP-1 mRNA, complete cds.	GGCGCAGCAT [C/T] CGCCTTTTCA	ນ ໝ	EH	H	H
G22u23	WIAF-10343	U86136	2877	Human telomerase-associated protein TP-1 mRNA, complete cds.	GCCCTCACC [G/A] TATCAGCCTT	ڻ س	4	ద	五
G22u24	WIAF-10344	U86136	3087	Human telomerase-associated 3087 protein TP-1 mRNA, complete cds.	TCAGGGCGCT [C/T] TGTGACAGAG	υ Σ	E	S	[24
G22u25	WIAF-10345	U86136	3662	Human telomerase-associated 3662 protein TP-1 mRNA, complete cds.	CAAGGTGGCA[C/T]CATTAGTCTT	υ Σ	E	<u>P</u> .	တ
G22u26	WIAF-10346	U86136	4762	Human telomerase-associated 4762 protein TP-1 mRNA, complete cds.	TTTCGAAGTT [C/T] CTTACCAACC	ນ ໝ	E-1	[Eq	<u>[24</u>
G22u27	WIAF-10351	U86136	1737	Human telomerase-associated 1737 protein TP-1 mRNA, complete cds.	CTCCAGCATG [G/C] GAAGTCGGTG	<u>ა</u>	υ		Ą
G22u28	WIAF-10352	U86136	3543	Human telomerase-associated 3543 protein TP-1 mRNA, complete cds.	ACAGTGCAAC[A/G]GCTGATGCTG	M	ט		м м
G22u29	WIAF-10353	U86136	4232	Human telomerase-associated 4232 protein TP-1 mRNA, complete cds.	GTCTGAGAGA[C/T]TCCGGACCCT	υ Σ	E	н	[E4
G22n30	WIAF-10354	U86136	4523	Human telomerase-associated 4523 protein TP-1 mRNA, complete cds.	GGAGGGCCCT [C/T] TGGAGCGCCC	ა ე	E-		ㅂ
G22u31	WIAF-10355	U86136	5333	Human telomerase-associated protein TP-1 mRNA, complete cds.	TGGTTGTCGG [G/T] TGCTGCAGAC	<u>ت</u>	E	>	니
G22u32	WIAF-10356	U86136	6208	Human telomerase-associated 6208 protein TP-1 mRNA, complete cds.	AGCTGCTGAC [G/A] CGGCCACACA	S	4	E	H

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G22u33	WIAF-10357	U86136	H 9 2777	Human telomerase-associated	TAGTGAGCCA [A/G] CACCACATCT	Æ	ט	T.	A
G22u34	WIAF-10360	U86136	3881 P	Human telomerase-associated 3881 protein TP-1 mRNA, complete cds.	CATCGAIGGC [G/A] CTGAIAGGIT	<u>ت</u>	A	4	E
G222111	WIAE-11700	M57230	1 t	IL6ST, interleukin 6 signal transducer (gp130, oncostatin M receptor)	TGAGTGGGAT [G/C]GTGGAAGGGA	ტ ჯ	Ų	U	px
Cucces	WTAF-11701	M57230		IL6ST, interleukin 6 signal transducer (gp130, oncostatin M receptor)	GTGGAAGGGA [A/G] ACACACTTGG	8	ບ	D2	ы
2222213	WTAF-11702	M57230	1 t t 577 x	ILGST, interleukin 6 signal transducer (gpl30, oncostatin M 677 receptor)	gagggaaga [a/g] aatgaggtgt	M	Ď	×	rx Px
222214	WIAF-11706	M57230	1616	IL6ST, interleukin 6 signal transducer (gp130, oncostatin M 1616 receptor)	aagaaatata [r/c]acttgagtgg	H E	υ	н	E
2222115	WTAF-11667	M57230	1444	<pre>IL6ST, interleukin 6 signal transducer (gp130, oncostatin M receptor)</pre>	TGATCGCTAT [C/G] TAGCAACCCT	D Z	v	н	>
G222u6	WIAF-11708	M57230	981	IL6ST, interleukin 6 signal transducer (gpl30, oncostatin M 981 receptor)	TCTTAAAATT [G/C] ACATGGACCA	<u>უ</u>	U	ъ	[IL]
G226u1	WIAF-11714	M85079	869	TGFBR2, transforming growth 869 factor, beta receptor II (70-80kD)	growth II (70-80kD) CACTGGGAGT[T/C]GCCATATCTG	E S	υ	۵	>
G226u2	WIAF-11715	M85079	1749	TGFBR2, transforming growth 1749 factor, beta receptor II (70-80kD)	wth (70-80kD) AGAITAIGAG[C/T]CTCCAITIGG	υ Σ	E	Д	w
G226u3	WIAF-11716	M85079	1601	TGFBR2, transforming gro factor, beta receptor II	wth (70-80kD) TGGGAACTGC[A/G]AGATACATGG	ν Δ		Æ	A
G226u4	WIAF-11721	M85079	1256	TGFBR2, transforming growth 1256 factor, beta receptor II (70-80kD)	growth II (70-80kD) TACTCCAGTT[C/G]CTGACGGCTG	υ Σ	<u></u> <u></u>	Ē	ы
G226u5	WIAF-11722	M85079	1502	TGFBR2, transforming growth 1502 factor, beta receptor II (70-80kD)	TCGTGAAGAA [C/T]GACCTAACCT	ນ ໝ	E⊣	z	Z
G226u6	WIAF-11671	M85079	888	TGFBR2, transforming growth 888 factor, beta receptor II (70-80kD)	growth II (70-80kD) TGTCATCATC[A/C]TCTTCTACTG	M	<u>υ</u>	<u>H</u>	1

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7119655	WTAF-11674	M85079	1425	1425 factor, beta receptor II (70-80kD)	II (70-80kD) CCTCCACAGT[G/A]ATCACACTCC	D E	Æ	Д	z
G22711	WIAF-10197	M86511	685	CD14 antigen		M	ro .	z	
G227u2	WIAF-10212	M86511	497		GAAGCCACAG [G/A] ACTTGCACTT	ت ع	A	U	м
				CUBN, cubilin (intrinsic factor-		E		2	2
G2278u1	WIAF-14117	AF034611	959	cobalamin receptor)	AGATAAATAA [T/C] GGCGGCTGTT	מ	ار	2	3
	01170	NE024611	781	CUBN, cubilin (intrinsic factor-	GGGTGGATGT [C/T] TTCACCCAAC	υ Σ	E	ß	ഥ
277 / 077	OTTE TOTA			CUBN, cubilin (intrinsic factor-				:	;
G2278u3	WIAF-14119	AF034611	641	cobalamin receptor)	CTGAGACGTA [C/T] GGACCCCAGT	<u>ာ</u>	=	н	X
				CUBN, cubilin (intrinsic factor-		<u>ر</u> خ	۲	_ρ	E
G2278u4	WIAF-14121	AF034611	1185		TGGTTATGGG [C/A] CAAATGGATG		£ _	4	-
				CUBN, cubilin (intrinsic factor-					2
G2278u5	WIAF-14133	AF034611	1532		TCTGGGTTAT [C/G] AAAACTGAAA	ر اد	١	-	Ξ.
A1187002	WTAF-14134	AF034611	2208	CUBN, cubilin (intrinsic factor-2208 cobalamin receptor)	GCCTTTCACT [C/T] ACACCAGGCA	υ Σ	H	н	H
250				IL10RA, interleukin 10 receptor,					í
G228u1	WIAF-10199	U00672	586	alpha	GCAAGGTGCC [G/A] GGAAACTTCA	S D	4	4	14
				IL10RA, interleukin 10 receptor,				<u></u>	Þ
G228u2	WIAF-10200	U00672	731	alpha	AGAGGAGTGC [A/G] TCTCCC TCAC	E E		-	> -
132280111	WIAF-13970	AJ001515	1747	1747 RYR3, ryanodine receptor 3	CAGGTATCTT [G/A] GAAGTTTTGC	ა ე	4	귀	ㅂ
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G2280u2	WIAF-13974	AJ001515	8593	8593 RYR3, ryanodine receptor 3	IAGAAGCCAI [I / C] GI CAGCAGI G	_		-	
G2282u1	WIAF-12694	D00726	263	FECH, ferrochelatase (protoporphyria)	ACATGGGAGG [C/T] CCTGAAACTC	8	E	Ŋ	_ U
22282112	WIAF-12695	D00726	514	FECH, ferrochelatase (protoporphyria)	TACTATATTG [G/A] ATTTCGGTAC	Σ Σ	4	ש	田
				ceo conronornhyrinogen oxidase					
G2285u1	WIAF-12688	D16611	673	(coproporphyria,	harderoporphyria) AGAAGACGCT[G/A]TCCATTTTCA	Σ Σ	A.	>	н
000000000000000000000000000000000000000	00 9 C L - B & F III	ח, אנת	783	CPO, coproporphyrinogen oxidase	ATCGTGGAGA [G/A] CGGCGGGGCA	S	4	[2]	四
2770277	NOOT TUIN			PrgER4 prostaglandin E receptor					
G2287u1	WIAF-12687	D28472	502	4 (subtype EP4)	GGGCCTCACG [C/T] TCTTTGCAGT	Σ	H	니	[14
C1787112	WTAF-12691	D28472	1309	PTGER4, prostaglandin E receptor 1309 4 (subtype EP4)	TGAAAATGGC [C/T] TTGGAGGCAG	×	H ن	니	[z _i
	70701-94TW	n28472	P7	PTGER4, prostaglandin E receptor 4 (subtype EP4)	AGGAGACGAC [C/I] IICIACACGC	<u>ω</u>	H U	H	₽
62287us	WIAF-IA/0/	7/1070							

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	C 7	0000	22.2	PTGER4, prostaglandin E receptor	GGTGTGCCTG [G/A] CATGGGCCTG	M G A	<u>ე</u>	Д
6228 / u4	WIRE-12/10	CAL AT1		DF1, stromal cell-derived factor		M G A	R F	Ж
G229611	WIAF-12727	25.055 0.0080		TB4R, leukotriene b4 receptor chemokine receptor-like 1)	CTATGTCTGC[G/C]GAGTCAGCAT	უ უ	r r	rk.
G2295u2	WIAF-12728	D89079	1248	LTB4R, leukotriene b4 receptor (chemokine receptor-like 1)	AGGGCACGGG [T/C] TCCGAGGCGT	υ Ε	g	U
G2295u3	WIAF-12753	D89079	1348	LTB4R, leukotriene b4 receptor 1348 (chemokine receptor-like 1)	CCTCACTGCC [T/G] CCAGCCCTCT	E E	S	A
G230u1	WIAF-10201	U31628	627	IL15RA, interleukin 15 receptor, alpha	acagccaaga [a/c] ctgggaactc	M A	z	E
G2300u1	WIAF-12735	J02959	102	102 LTA4H, leukotriene A4 hydrolase	ACCTGCACCT [G/T] CGCTGCAGCG	S G T	ы	ы
G2300u2	WIAF-12738	J02959	1380	LTA4H, leukotriene A4 hydrolase	CCTGGCTCTA [C/T] TCTCCTGGAC	S C	ъ	Þı
G2302u1	WIAF-12741	J03037	627	627 CA2, carbonic anhydrase II	TCCTGAATCC [C/T] TGGATTACTG	S C	ᄓ	ы
G2302u2	WIAF-12742	J03037	819	819 CA2, carbonic anhydrase II	GCCACTGAAG [A/G] ACAGGCAAAT	M A B	Z	Д
G2303u1	WIAF-12751	J03571	304	ALOX5, arachidonate 5- lipoxygenase	CGCTGAAGAC [G/A] CCCCACGGGG	ა დ	H	E-1
G2303u2	WIAF-12752	J03571	794	ALOX5, arachidonate 5- lipoxygenase	AGAGCTGCCC [G/A] AGAAGCTCCC	M D	[2]	X
G2304u1	WIAF-12772	J03575	840	PDHA1, pyruvate dehydrogenase (lipoamide) alpha 1	TCCGAGAGGC [A/G] ACAAGGTTTG	ନ ୟ ହ	K.	Ą
G2304u2	WIAF-12779	303575	1044	PDHA1, pyruvate dehydrogenase (lipoamide) alpha 1	CCAGTGTGGA [A/C] GAACTAAAGG	M A	四	Д
G2305u1	WIAF-12763	J03576	456	PDHB, pyruvate dehydrogenase (lipoamide) beta	TCTTCAGGGG [A/G] CCCAATGGTG	ر الا		υ
32305112	WIAF-12764	J03576	650	PDHB, pyruvate dehydrogenase (lipoamide) beta	GITCCITITG [A/C] ALTICICCCG	M A	ы	4
G231u1	WIAF-10202	U32324	734	ILLIRA, interleukin 11 receptor, 734 alpha	CCAGGGCCTG [C/T]GGGTACAGTC	M C	ద	⋈

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1,10,100	WI2H-10762	30501.	3726	rting, alpha 2 (+)	TCAAGAACCA [C/T] ACAGAGATCG	S C	田	田
02312u1	WTAR-12760	105200	6141	odine receptor 1	TGCAATTCAA [A/G]GATGGTACAG	S	ى بى	
9401045	WTAF-12767	705200		nodine receptor 1	CGGCGCAGAC [A/G] ACACTGGTGG	8	Ð	E
G231313	WIAF-12768	305200	3084	nodine receptor 1	ATGGGCACAA [C/T] GTGTGGGCCC	ა ე	Z H	Z
G2313114	WIAF-12777	705200	1 5667	RVR1, ryanodine receptor 1 (skeletal)	GCATCTTTGG [C/T] GATGAGGATG	ე დ	E	n D
22313115	WIAF-12780	002500	0099	RYR1, ryanodine receptor 1 (skeletal)	GCTCGCTGCT [C/T] ATCGTGCAGA	ე დ	E	ㅁ
G2313u6	WIAF-12781	J05200	7191	RYR1, ryanodine receptor 1 (skeletal)	AGCCTGAGTG [C/T] TTCGGACCCG	υ 8	EH	U U
G2313u7	WIAF-12782	J05200	7602	RYR1, ryanodine receptor 1 (skeletal)	ACCACAAGGC [G/A] TCCATGGTGC	හ ව	Æ	A
G2313u8	WIAF-12784	J05200	9288	RYR1, ryanodine receptor 1 (skeletal)	CAGACGCCCC [A/G] GCTGTGGTCA	S A	ъ	<u>д</u>
G2313u9	WIAF-12786	J05200	13690	RYR1, ryanodine receptor 1 13690 (skeletal)	TCCAAAGAAG [G/A] AGGAAGCTGG	<u>ნ</u>	A	四
G2313u10	WIAF-12789	J05200	3147	RVR1, ryanodine receptor 1 (skeletal)	ACATCCCAGC [G/A] CGCCGAAACC	ა ე	A	A A
G2314u1	WIAF-12771	J05272	1920	IMPDH1, IMP (inosine monophosphate) dehydrogenase 1	TGAAGATCGC [A/G] CAGGGTGTCT	∀	ъ	4 4
G2319u1	WIAF-12814	K03191	651	CYP1A1, cytochrome P450, subfamily I (aromatic compound- inducible), polypeptide 1	CCCCTACAGG [T/C] ATGTGGTGGT	E E	υ	<u>н</u> х
G232u1	WIAF-11657	U58917	1490	Homo sapiens IL-17 receptor mRNA, complete cds.	TGAACATGAT [C/T] CTCCCGGACT	ა ა	H	Н
G232u2	WIAF-11677	U58917	1293	Homo sapiens IL-17 receptor mRNA, complete cds.	GCAGGCCATC [T/C] CGGAGGCAGG	E N	υ	ري ب
G232u3	WIAF-11658	U58917	1132	Homo sapiens IL-17 receptor mRNA, complete cds.	GGCCTGCCTG [C/T] GGCTGACCTG	Ω Z	E	D A
G232u4	WIAF-11679	U58917	902	Homo sapiens IL-17 receptor mRNA, 905 complete cds.	GCAGCTGCCT[C/T]AATGACTGCC	S C	H	<u>1</u>

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(3232u5	WIAF-11682	U58917	1 1794 c	Homo sapiens IL-17 receptor mRNA, 1794 complete cds.	GTTCGAATGT [G/T] AGAACCTCTA	Ö	H	ш	*
	C () () () () () () () () () (7 10 0 0 1	47 74	Homo sapiens IL-17 receptor mRNA,	TGACCAGTTT [T/C] CCGCACATGG	<u>٦</u> د	ŭ	ĺΉ	[II.
623247	OCCUT - JAIN	1			M TOPERATION TO STRANGEST	T	υ	Σ	<u></u>
G2322u1	WIAF-12853	L01406	1285 GCGR.	1316 MOIMONE TECEPOOL 1285 GCGR. glucagon receptor			บ	ж	PK
G2329u1	WIAF-12850	L22214	713 2	+		E .	ŭ	Н	н
G2329u2	WIAF-12851	L22214	716	716 ADORA1, adenosine Al receptor	TGGCAATTGC[T/G]GTGGACCGCT	S E	<u></u>	Ą	Æ
G2335a1	WIAF-12136	132961	265	ABAT, 4-aminobutyrate aminotransferase	CCTAGATCTC[A/G]GGAGTTAATG	M A	_O	α	ద
G2335a2	WIAF-12137	L32961	407	ABAT, 4-aminobutyrate	TCTCCTCTGT [T/C] CCCATAGGTT	S	ŭ	Δ	>
(42335)13	WIAF-12838	132961	365	ABAT, 4-aminobutyrate aminotransferase	TTGATGTGGA[C/T]GGCAACCGAA	ა ე	E	Д	Д
7132500	WTAF-12839	1,32961	583	ABAT, 4-aminobutyrate aminotransferase	ATCACCATGG [C/T] CTGCGGCTCC	υ Σ	E	Ø	Λ
42335115	WIAF-12841	L32961	1082	ABAT, 4-aminobutyrate aminotransferase	TGGACGAGGT [C/A] CAGACCGGAG	υ 0	4	>	Λ
33546	WTAF-12852	1,32961	227	ABAT, 4-aminobutyrate aminotransferase	ATTATGATGG [G/A] CCTCTGATGA	დ დ	A	Ü	ro l
				ALDH5Al, aldehyde dehydrogenase 5					
G2337u1	WIAF-13577	L34820	149	ialdehyde dehyd	TGTTCTCGAA [A/G] GAATGCCAAG	M C	ტ [X F	요 누
G2342a1	WIAF-12138	M12530	1602 TF,	TF, transferrin		1.) [-	<u> </u>	S
G2342a2	WIAF-12139	M12530	AIA AIA 234 Geh	la S	TGGCCAGGTA [T/C] GGTGTGAAGC	Ω [-1	٥	>1	Y
62346u1	WTAF-12830	M13928	529	ALAD, aminolevulinate, delta-, 529 dehydratase	TGAGGTGGCA [T/C] TGGCGTATGC	S	ŭ	ᆈ	ц
G2346u3	WIAF-12843	M13928	480	ALAD, aminolevulinate, delta-, 480 dehydratase	TGAGTGAAAA [C/T] GGAGCATTCC	S	H		Z
G2348u1	WIAF-12835	M14016	621	UROD, uroporphyrinogen 621 decarboxylase	CTCTGGTCCC [A/G] TATCTGGTAG	Ω Æ	<u></u>	Δı	Д

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(2) 2 E 11 1	WTAF-11678	1183171	100	SCYA22, small inducible cytokine 100 subfamily A (Cys-Cys), member 22	CAGGCCCCTA[C/T]GGCGCCAACA	დ დ	E→		X
100000	9 1 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	M37435	596	Н	GACAAGGACT [G/T] GAATATTTTC	Ŭ Œ	H	Z	ᄓ
2889828	WIAF-13225	M37435	498	y stimulating factor 1	AAGAGCATGA [C/T] AAGGCCTGCG	ပ တ	Ę-i	Д	
G2363a3	WIAF-13226	M37435	712	CSF1, colony stimulating factor 1	CAGTGACCCG [G/T] CCTCTGTCTC	IJ E	H	A	w
G2369u1	WIAF-12854	M30773	857	PPD3R1, protein phosphatase 3 (formerly 2B), regulatory subunit B (19kD), alpha isoform (calcineurin B, type I)	TTGATTTGGA [C/T] AATTCTGGTT	ω υ	E	Δ	Д
9236902	WIAF-12855	M30773	1274	PPP3R1, protein phosphatase 3 (formerly 2B), regulatory subunit B (19kD), alpha isoform 1274 (calcineurin B, type I)	ATGTGTGACT [C/T] TTATCAGAGA	υ I	E		
1,1720	WTAF-11662	U86358	311	SCYA25, small inducible cytokine subfamily A (Cys-Cys), member 25	CACCACAACA [T/C] GCAGACCTTC	E	<u>υ</u>	Σ	<u> </u>
	0001	α υ υ	134	SCYA25, small inducible cytokine 134 subfamily A (Cvs-Cvs), member 25	GTGCTCCGGC [G/A] CGCCTGGACT	M	A	ρ;	田
G237u2 G237u3	WIAK-II68U	U86358	133	SCYA25, small inducible cytokine subfamily A (Cys-Cys), member 25	TGTGCTCCGG [C/T] GCGCCTGGAC	D E	Et	ద	υ
7.1.7 7.1.7	WTAF-11661	U86358	302		GCAAAGCTCC [A/G] CCACAACATG	M 4	r c		ద
	MT 2 11 5 6 3	11 87 87 87 87 87	378	SCYA25, small inducible cytokine subfamily A (Cys-Cys), member 25	AGTTATCA [A/G] TCCAAGTTTA	ν 4	<u> </u>	Ø	<u> </u>
G23 7311	WIAF-12870	M36035	500	BZRP, benzodiazapine receptor (peripheral)	GCTGGCCTTC [G/A] CGACCACACT	<u>υ</u>	A	- A	<u></u>
1,19250	WTAR-13025	M57414	979	979 TACR2, tachykinin receptor 2	CTGCTGCCCA [T/C] GGGTCACACC	Ε	Ü		ద
G238u1	WIAF-10177	X01394	235	lan ti	GCTCCAGGCG [G/T] TGCTTGTTCC	8	G H	<u> </u>	

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22381111	WIAF-12894	M59941	730	CSF2RB, colony stimulating factor 2 receptor, beta, low-affinity (granulocyte-macrophage)	CAGAGGTTTG [C/T] TGGGACTCCC	υ	D E	Ü
G2381u2		M59941	1306	ing factor finity	GGATCTGGAG [C/T] GAGTGGAGTG	U m	E .	<u>လ</u>
G2381u3	WIAF-12900	M59941	1972	ing factor finity	CGATGGGACC [G/A] GGACAGGCCG	ω v	A	<u>д</u> ,
G2381u4	WIAF-12901	M59941	1982	CSF2RB, colony stimulating factor 2 receptor, beta, low-affinity (granulocyte-macrophage)	GGGACAGGCC [G/A] TGGAAGTGGA	<u>უ</u>	Æ	M D
20 20 20 20 20 20 20 20 20 20 20 20 20 2	WIAF-12942	M59941	773	CSF2RB, colony stimulating factor 2 receptor, beta, low-affinity (granulocyte-macrophage)	CCAGAACCTG [G/C]AGTGCTTCTT	<u>უ</u>	υ	O E
0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.0	WIAF-12946	M59941	2458	CSF2RB, colony stimulating factor 2 receptor, beta, low-affinity (granulocyte-macrophage)	CCCCACAGCC [C/A] GAGGGCCTCC	ပ အ	A	<u>а</u>
G2384u1	WIAF-12908	M61831	1000	AHCY, S-adenosylhomocysteine	GCCGTGGAGA [A/C] GGTGAACATC	Æ	υ	R F
G2387u1	WIAF-12910	M63967	2585	2585 ALDH5, aldehyde dehydrogenase 5	CTGCTGAACC[T/G]CCTGGCAGAC	E E	_O	디
G2387u2	WIAF-12911	M63967	2996	2996 ALDH5, aldehyde dehydrogenase 5	TATGGCCCAA [C/G] AGCAGGTGCG	υ Σ	ъ	H H
G2387u3	WIAF-12954	M63967	2522	2522 ALDH5, aldehyde dehydrogenase 5	GCCCGGGAAG [C/T] CTTCCGCCTG	υ Σ	H	N A
G2387u4	WIAF-12955	M63967	2448	2448 ALDH5, aldehyde dehydrogenase 5	ACCCTACCAC[C/T]GGGGAGGTCA	ე ც	E	T

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42387115	WIAF-12956	M63967	2460	2460 ALDH5, aldehyde dehydrogenase 5	GGGAGGTCAT [C/T] GGGCACGTGG	ບ ໝ	터	н	н
G2387u6	WIAF-12957	M63967	2991	2991 ALDH5, aldehyde dehydrogenase 5	CGGGGTATGG [C/T] CCAACAGCAG	ر د	Ŀ	υ	_D
G2387u7	WIAF-12958	M63967	3022	3022 ALDHS, aldehyde dehydrogenase 5	CGCCCAGCAC [A/G] TGGATGTTGA	Æ	_ ტ	Σ_	Δ
G2387u8	WIAF-12959	M63967	2943	ALDH5, aldehyde dehydrogenase 5	CCCTCATCAA [G/C] GAGGCAGGCT	₽	υ	×	Z
G2388u1	WIAF-12888	M64590	88 10	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 588 system protein P)	TGCCACAGAC [G/A] ATTTTGCGGA	ري 2		EH	Ę-I
G2388u2	WIAF-12889	M64590	651	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 651 system protein P)	ACCAGCCTGA [G/A] GTGTCTCAGG	დ დ	A.	匝	E
G2388u3	WIAF-12890	M64590	869	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 898 system protein P)	CAGACCATGG [T/C] GTGTGACATC	E E	<u></u> <u></u>	>	A
G2388u4	WIAF-12891	M64590	557	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 557 system protein P)	TATATTGGCA [T/C] GGGCTATTAT	E E	Ü	Σ	E
22.28.83.1.5	MTAR-12938	M64590	587	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 587 system protein P)	GTGCCACAGA[C/G]GATTTTGCGG	υ Σ	<u> </u>	EH	~ ~
G2388u6	WIAF-12939	M64590	518	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage 518 system protein P)	CTGCATGCCA [T/C] TTCAAGCAAA	Σ	D H		타

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G2388117	WIAF-12940	M64590	810	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage slo system protein P)	GGAAATTTCT [C/T] GTTGATCCCC	ر ن ا	<u>ы</u>	д
G2388u8	WIAF-12941	M64590	1481	SLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage system protein P)	CATTGTGGCT [G/A] CTCAGTGAAG	ع ب ع	U	>
G2388u9	WIAF-12947	M64590	1841	dehydrogenase 1g, glycine glycine cleavage P)	AAACTGAACA [G/A] TTCGTCTGAA	ى ب	ري د	
23381110	WIAR-12948	M64590	2325	GLDC, glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage	GACAGGICTA [C/T] CTAGACGGGG	ე დ		<u>></u>
	# H#	М. А. А. В.	23,62	dehydrogenase 1g; glycine glycine cleavage p)	GGTGGGAATC [T/A] GTCGCCCTGG	El E		<u>N</u>
62388ull	WIAE-12949	M64590	. 3220	dehydrogenase ng; glycine glycine cleavage P)	ttagtcctct [c/g] tccctaagtt	ن ا	ტ	1
G2391u1	WIAF-12998	M69238	623	ARNT, aryl hydrocarbon receptor 623 nuclear translocator	TGGTGTATGT [G/C] TCTGACTCCG	<u>დ</u>	Ü	<u>></u>
G2391u2	WIAF-13002	M69238	1072	ARNT, aryl hydrocarbon receptor 1072 nuclear translocator	TGCCTAGTGG [C/T] CATTGGCAGA	υ Σ	E E	A
62391113	WIAF-13021	M69238	996	ARNT, aryl hydrocarbon receptor 966 nuclear translocator	ACCTCACTTC [G/A] TGGTGGTCCA	o M	Æ	∑ >
G2394u1	WIAF-13003	M73747	2061	TSHR, thyroid stimulating hormone 2061 receptor	TTGCTGGTAC [T/A] CTTCTATCCA	E E	Æ	H I

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C1178672	WTAF-13004	M73747	2248	TSHR, thyroid stimulating hormone 2248 receptor	TTACCCACGA [C/G] ATGAGGCAGG	ν υ	G	ū	闰
G2396u1	WIAF-12995	M74542	1027	1027 ALDH3, aldehyde dehydrogenase 3	CCCCCAGTCC [C/G] CGGTGATGCA	<u>ν</u>	ט	ф	A
G2396u2	WIAF-13019	M74542	1295	1295 ALDH3, aldehyde dehydrogenase 3	GGCAAGAAGA [G/A] CTTCGAGACT M	<u>ح</u>	Æ	က	z
G2403u1	WIAF-13583	M83670	280	CA4, carbonic anhydrase IV	TACGATAAGA [A/T] GCAAACGTGG	₹	H	×	Σ
G2409u1	WIAF-10010	HT2156	1268	1268 AGTR1, angiotensin receptor 1	CCACTCAAAC[C/T]TTTCAACAAA	υ Σ:	E	ᄓ	ĹΉ
G2411u1	WIAF-13541	M97759	210	210 ADORA2B, adenosine A2b receptor	TGGCGGGCAA [C/T] GTGCTGGTGT	ა ე	E	z	z
G2422u1	WIAF-14077	890469	375	POR, P450 (cytochrome) oxidoreductase	GCAGCCTGCC [A/G] GAGATCGACA	SA	ಅ	Д	Д
G2422u2	WIAF-14078	890469	852	POR, P450 (cytochrome) oxidoreductase	TCCTGGCTGC [A/G] GTCACCACCA	∀	೮	Æ	Æ
G2422u3	WIAF-14082	890469	1496	POR, P450 (cytochrome) 1496 oxidoreductase	AAGGAGCCTG [T/C] CGGGGAGAAC	E	υ	>	ø
G2422u4	WIAF-14099	890469	1443	POR, P450 (cytochrome) oxidoreductase	AGACCAAGGC [C/T] GGCCGCATCA	Ω Ω	H	4	A
G2422u5	WIAF-14100	890469	1704	POR, P450 (cytochrome) oxidoreductase	GCCGCCGCTC [G/A] GATGAGGACT	ა დ	Æ	တ	ß
G2427u1	WIAE-14079	U07919	1369	ALDH6, aldehyde dehydrogenase 6	ACTATGGACT [C/T] ACAGCAGCCG	S S	H		ㅁ
G2427u2	WIAF-14096	007919	1347	1347 ALDH6, aldehyde dehydrogenase 6	ATAAAAAGAG[C/T]GAATAGCACC	υ Σ	H	A	>
G243u1	WIAF-11684	X57522	926	TAP1, transporter 1, ABC (ATP 926 binding cassette)	ATAGCCAGTG [C/G] AGTGCTGGAG	υ Σ	U	<<	Ŋ
G243112	WIAF-11685	X57522	627	TAP1, transporter 1, ABC (ATP 627 binding cassette)	ACCCTACCGC [C/T] TTCGTTGTCA	S C	H	< 4	∢
G243u3	WIAF-11686	X57522	538	TAP1, transporter 1, ABC (ATP 538 binding cassette)	CCTGCCGGGA [C/G] TTGCCTTGTT	υ Σ	υ	ㅁ	⊳
G243u4	WIAF-11687	X57522	798	TAP1, transporter 1, ABC (ATP binding cassette)	TGGTGGTCCT [C/G] TCCTCTTG	ა ე		ㅁ	ы
G243u5	WIAF-11689	X57522	1465	TAP1, transporter 1, ABC (ATP 1465 binding cassette)	TAGTATTTCA [G/T] GTATGCTGCT	Σ Σ	E+	ტ	υ
G243u6	WIAF-11690	X57522	177	TAP1, transporter 1, ABC (ATP 177 binding cassette)	AGAGTCCCAG [A/G] CCCGGCCGGG	S A	Ŋ	ద	p4
G243u7	WIAF-11693	X57522	1067	TAP1, transporter 1, ABC (ATP 1067 binding cassette)	AACATCATGT [C/T] TCGGGTAACA	Σ Σ	H	တ	<u>[</u>
G243u8	WIAF-11665	X57522	1207	TAP1, transporter 1, ABC (ATP 1207 binding cassette)	GGTCACCCTG [A/G] TCACCCTGCC	M	Ŋ	. н	>

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C2443U3 WIAF-11664 X57522 1757 Dinding cassette) ABG (APP Dinding cassette) ABG (APP Dinding cassette) ABG (APP ABF-10174 X60592 239 Peceptor superfamily, member 5 SIC12A2, solute carrier family (and and and and and and and and and and						-	L	_	Ŀ
MIAF-10174 X60592 239 receptor s	 	X57522	1757	r 1, ABC (ATP	CCAAACCGCC [C/T] AGATGTCTTA	Σ.	[-i	д	ы
SIC12A2, (sodium/poorte 1355 transporte 1360 transporte 1355 transporte 1355 transporte 1355 transporte 1356 transporte 13		X60592	239	ω	CTTGCGGTGA [A/G]AGCGAATTCC	S A	U	Щ	田
SICC12A2, (sodium/pc (sodium/pc (sodium/pc (sodium/pc (sodium/pc (sodium/pc (sodium/pc (subfamily (subfa	 	U30246	1355	SLC12A2, solute carrier family 12 (sodium/potassium/chloride transporters), member 2	IGCTTAAGGA [A/G] CATTCCATAC	ر ا	Ö	田	闰
CYP2J2, cytochrome P450, subfamily IIJ (arachidonic acis subfamily IIJ (arachidonic acis subfamily IIJ (arachidonic acis subfamily III (arachidonic acis subfamily III (arachidonic acis subfamily III) (arachidonic acis subfamily I	AF-13714	U30246	2691	SLC12A2, solute carrier family 12 (sodium/potassium/chloride transporters), member 2	AGCCAAATAT [C/G]AGCGATGGCT	Σ	ro e		四
CYP2J2, cytochrome P450, subfamily IIJ (arachidonic acia subfamily IIJ (arachidonic acia epoxygenase) polypeptide 2	AF-14004	U37143	1456		CTGAAGTTTA [G/A] AATGGGTATC	Σ Σ	4	ద	
CYP2J2, cytochrome P450, subfamily IIJ (arachidonic aci) WIAF-14033 U37143 1502 epoxygenase) polypeptide 2 WIAF-14065 U37519 771 ALDH3, aldehyde dehydrogenase WIAF-14114 U38178 236 phosphodiesterase WIAF-14115 U38178 849 phosphodiesterase WIAF-1412 U38178 1655 phosphodiesterase WIAF-14241 X95520 941 phosphodiesterase WIAF-14241 WIAF	AF-14032	U37143	376		tttaagaaa [a/g] tggattgaft	Æ	<u> </u>	z	Ω O
WIAF-14065 U37519 771 ALDH3, aldehyde dehydrogenase WIAF-14066 U37519 1698 ALDH3, aldehyde dehydrogenase WIAF-14114 U38178 236 phosphodiesterase WIAF-14115 U38178 249 phosphodiesterase WIAF-14122 U38178 21,31-cyclic nucleotide WIAF-14122 U38178 1655 phosphodiesterase WIAF-14241 X95520 941 phosphodiesterase	AF-14033	U37143	1502	CYP2J2, cytochrome P450, subfamily IIJ (arachidonic acid epoxygenase) polypeptide 2	TCTGCGCTGT [T/A] CCTCAGGTGT	ω H	4	>	> >
WIAF-14066 U37519 1698 ALDH3, aldehyde dehydrogenase WIAF-14114 U38178 236 phosphodiesterase WIAF-14115 U38178 849 phosphodiesterase CNP, 2',3'-cyclic nucleotide WIAF-1412 U38178 849 phosphodiesterase WIAF-1412 U38178 CNP, 2',3'-cyclic nucleotide WIAF-1421 U38178 1655 phosphodiesterase WIAF-14241 X95520 941 phosphodiesterase	AF-14065	U37519	771	ALDH3, aldehyde dehydrogenase 3	CCCGCAGGGA [A/G] TTGCGTGGTG	M	A G	Z	വ
CNP, 2',3'-cyclic nucleotide WIAF-14114 U38178 236 phosphodiesterase CNP, 2',3'-cyclic nucleotide WIAF-14115 U38178 849 phosphodiesterase CNP, 2',3'-cyclic nucleotide WIAF-14122 U38178 1655 phosphodiesterase CNP, 2',3'-cyclic nucleotide CNP, 2',3'-cyc	AF-14066	U37519	1698	, aldehyde	AAGGAGATCC [G/A] CTACCCACCC	Σ	ত ত	ద	田
CNP, 2',3'-cyclic nucleotide WIAF-14115 U38178 849 phosphodiesterase CNP, 2',3'-cyclic nucleotide WIAF-14122 U38178 1655 phosphodiesterase CNP, 2',3'-cyclic nucleotide WIAF-14241 X95520 941 phosphodiesterase P41 Phosphodiesterase P42 Phosphodiesterase P42 Phosphodiesterase P43 Phosphodiesterase P43 P44 Phosphodiesterase P43 P44 P4	AF-14114	U38178	236	r)	TGCCGGGCGC [G/A] CCTCTCGCTG	Σ	ج ن	. R	ш
WIAF-14122 U38178 1655 phosphodiesterase CNP, 2',3'-cyclic nucleotide CNP, 2',3'-cyclic nucleotide WIAF-14241 X95520 941 phosphodiesterase	AF-14115	U38178	849	CNP, 2',3'-cyclic nucleotide 3' phosphodiesterase	GTGCCGCCGA [A/G] GAAAAAGTGC	w	<u>ن</u> 4	四	<u> </u>
CNP, 2',3'-cyclic nucleotide WIAF-14241 X95520 941 phosphodiesterase	.AF-14122	U38178	1655	CNP, 2',3'-cyclic nucleotide 3' phosphodiesterase	GTTATCTTGC[A/T]GAGATCTCTG	Σ	A		J Ø
	 [AF-14241	X95520	941	CNP, 2',3'-cyclic nucleotide 3' phosphodiesterase	TGCAAAATAT [T/C] CAGGAGACCG	٥٠	- C		۷۰

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G2445u5	WIAF-14242	X95520	1057	CMP, 2',3'-cyclic nucleotide 3'	TGGAGTTGAT [C/T] TTTCAGTGCT	U U	٥٠	٥٠
G2445u6	WIAF-14243	X95520	1583	CNP, 2',3'-cyclic nucleotide 3' phosphodiesterase	TCTACTGGCT [C/G] TCTAACTAAT	D U	٥٠	٥.
G2448u1	WIAF-13973	U46689	1895	ALDH10, aldehyde dehydrogenase 10 (fatty aldehyde dehydrogenase)	TTGTCAAGGC [A/T] GAATATTACT	A	T A	4
G2457u1	WIAF-13898	090277	1304	GRIN2A, glutamate receptor, ionotropic, N-methyl D-aspartate 2A	GGTCCCGATG [C/T] ACACCTTGCA	υ	표	>1
G2457u2	WIAF-13899	090277	GR. 1934 2A	IN2A, glutamate receptor, notropic, N-methyl D-aspartate	AAGAAGTAAT [G/T] GCACCGTCTC	හ	₽ -	υ
G2457u3	WIAF-13900	U90277	GR. 101 2230 2A	GRIN2A, glutamate receptor, ionotropic, N-methyl D-aspartate 2A	TCGCTGTCAT [A/G] TTCCTGGCTA M	A	D D	Σ
G2457u4	WIAF-13902	U90277	2916	GRIN2A, glutamate receptor, ionotropic, N-methyl D-aspartate 2A	GGCATCTACA [G/A] CTGCATTCAT M	Ö	8	
G2457u5	WIAF-13903	U90277	3251	GRIN2A, glutamate receptor, ionotropic, N-methyl D-aspartate 2A	CTATGTATTC[C/T]AGGGACAACA	U	O E	*
G2457116	WIAF-13917	U90277	2756	GRINZA, glutamate receptor, ionotropic, N-methyl D-aspartate 2A	GGACATTGAC [A/G]ACATGGCGGG	A	<u>ත</u>	
G2468u1	WIAF-13642	X04011	1017	CYBB, cytochrome b-245, beta polypeptide (chronic granulomatous 1017 disease)	AGGTGTCCAA [G/A] CTGGAGTGGC	ტ	A A	M
G2473u1	WIAF-13670	06690X	1417	ICAM1, intercellular adhesion molecule 1 (CD54), human 1417 rhinovirus receptor	GGTCACCGC [G/A] AGGTGACCGT	υ Σ	A	<u> </u>
G2473u2	WIAF-13695	06690X	179	ICAM1, intercellular adhesion molecule 1 (CD54), human 179 rhinovirus receptor	GACCAGCCCA [A/T] GTTGTTGGGC	Σ Z	EH	Σ Σ
G2480u1	WIAF-14148	X55330	800	800 AGA, aspartylglucosaminidase	TIGGCAIGGI[I/G]GIAAICCAIA	<u>ا</u>	r _o	Δ Δ
G2480u2	WIAF-14149	X55330	852	852 AGA, aspartylglucosaminidase	AAATGGTATA [A/T] AATTCAAAAT	N A	H	* ×

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6::00	MTAR-14158	X55330	616 AGA	aspartvlqlucosaminidase	TTATCTACCA [G/C] TGCTTCTCAA	r E	ט	Ø.	₽
G2485u1	WIAF-13612	X59543	2301	se M1	ATTGATCAAA [G/A] CCAATCTTTG	<u>ი</u>	4	က	z
(22485112	WIAF-13613	X59543	2410	RRM1, ribonucleotide reductase M1 polypeptide	ATTTAAGGAC [G/A] AGACCAGCAG	ა ი	Æ	Eн	EH
G2485113	WIAF-13651	X59543	548	nucleotide reductase M1	CAAGTCAACA [T/C]TGGATATTGT	S F	U	н	ы
G248 F114	WTAF-13652	X59543	199	nucleotide reductase M1	TGCATGTGAT [C/T] AAGCGAGATG	ຮ	∺	н	н
G2485115	WTAF-13653	X59543	1037	RRM1, ribonucleotide reductase M1 polypeptide	CAACACAGCT [C/A] GATATGTGGA	ນ ຜ	Æ	ద	24
2248 End	WT&R-13660	X59543	1955	RRM1, ribonucleotide reductase M1 polypeptide	GAAGATTGCA [A/C]AGTATGGTAT	M	ŭ	×	Q
22485117	WTAF-13877	X59543	860	RRM1, ribonucleotide reductase M1 polypeptide	GAGTATGAAA [G/C]ATGACAGCAT	M G	υ	Ω	н
G2486u1	WIAF-14075	X59618	543	RRM2, ribonucleotide reductase M2 polypeptide	TCAGCACTGG [G/C] AATCCCTGAA	υ Σ	บ	四	Q
G2486u2	WIAF-14076	X59618	189	RRM2, ribonucleotide reductase M2 polypeptide	TCGCTGCGCC [T/G] CCACTATGCT	E I	<u></u> છ	- 1	1
G2486u3	WIAF-14092	X59618	524	RRM2, ribonucleotide reductase M2 polypeptide	TTGACCTCTC[C/G]AAGGACATTC	8	ರ	ß	S
G2488u1	WIAF-13585	X63563	1633	POLR2B, polymerase (RNA) II (DNA directed) polypeptide B (140kD)	CCTTGATGGC [G/A] TATATTTCAG	დ ტ	A	A	A
G2488u2	WIAF-13586	X63563	2452	POLR2B, polymerase (RNA) II (DNA directed) polypeptide B (140kD)	CTGTAGACCG [C/T] GGCTTCTTCA	ა ე	E-4	ద	ద
G2488u3	WIAF-13587	X63563	2740	POLR2B, polymerase (RNA) II (DNA 2740 directed) polypeptide B (140kD)	TCAGAACTAG [T/C] GAGACGGGCA	Ω H	U .	Ø	ß
G2488u4	WIAF-13602	X63563	1411	POLR2B, polymerase (RNA) II (DNA directed) polypeptide B (140kD)	GGGGTGATCA [A/G] AAGAAAGCTC	ر ا	<u> </u>	O)	Q
G2488u5	WIAF-13603	X63563	2386	POLRZB, polymerase (RNA) II (DNA directed) polypeptide B (140kD)	CAAITGIGGC [C/I]AITGCAICAI	<u>ي</u> 2	E	&	Æ
G2489u1	WIAF-14181	X63564	1346	POLR2A, polymerase (RNA) II (DNA 1346 directed) polypeptide A (220kD)	TGGTGGACAA [T/C] GAGCTGCCTG	ν Γ	D H	Z	z

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G2489u2	WIAF-14236	X63564	1847	1847 directed) polypeptide A (220kD)	TGAATCTTAG[C/T]GTGACAACTC	υ 0	£-1	۸.	٥.
G2489u3	WIAF-14237	X63564	2678	POLR2A, polymerase (RNA) II (DNA 2678 directed) polypeptide A (220kD)	CTGAATACAA [C/T] AACTTCAAGT	υ n	Ð	۵.	٥.
G2489u4	WIAF-14238	X63564	3059	POLRZA, polymerase (RNA) II (DNA 3059 directed) polypeptide A (220kD)	AGCTGCGCTA [C/T] GGCGAAGACG		₽	٠٠	٥.
G2489u5	WIAF-14239	X63564	3827	POLR2A, polymerase (RNA) II (DNA 3827 directed) polypeptide A (220kD)	TGGGCCAGTC[C/T]GCTCGAGATG	U C	E	٥.	Ç.
G2489u6	WIAF-14240	X63564	3992	POLRZA, polymerase (RNA) II (DNA directed) polypeptide A (220kD)	TGCCTGACTT [T/C] GATGTGGCCC	E1	ŭ	٥٠	٠.
G2489u7	WIAF-14245	X63564	3938	POLR2A, polymerase (RNA) II (DNA 3938 directed) polypeptide A (220kD)	CCCAGAGCAC [G/A] GTGGTGGCAG	<u>ن</u> .ه	Æ	٥.	٠.
G250u1	WIAF-11696	HT0155	1113	IL3RA, interleukin 3 receptor, alpha (low affinity)	CTGTGTCTTC[G/C]TGATCTGCAG M	υ Σ	υ .	<b>&gt;</b>	Д
G251111	WIAF-11666	HT0240	179	179 interleukin 1 beta convertase	TGGATAAGAC[C/T]CGAGCTTTGA	S S	E	EH	E
G251u2	WIAF-11694	HT0240	973	interleukin 1 beta convertase	GATGCTATTA [A/G] GAAAGCCCAC	ď	r to	×	ద
G251u3	WIAF-11695	HT0240	783	783 interleukin 1 beta convertase	CCCAGATATA[C/T]TACAACTCAA	<u>ე</u>	H	ᆸ	H
G2513u1	WIAF-13736	HT27365	1721	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	AACTATCTAT [G/A] AAAGCCAAA	<u>ა</u>	4	Σ	н
G2513u2	WIAF-13737	HT27365	1741	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	AACTATTGGG [A/T] AATGTGTTCA M	Z Σ	F	ы	>
G2513u3	WIAF-13738	HT27365	1697	PLCB3, phospholipase C, beta 3 1697 (phosphatidylinositol-specific)	AATCTGTTCA [A/G] TACAGGGATT	<u>م</u>	_O	Ø	O.

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WI	WIAF-13739	HT27365	1908	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	CTGTCAGATT [G/A] TAGCAATGAA	_O	A	<u>н</u>
3	WIAF-13740	HT27365	2172	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	TATAGAGATA[C/T]ACGGAATTCC M	υ	H	H H
	WIAE-13744	HT27365	3019	PLCB3, phospholipase C, beta 3 3019 (phosphatidylinositol-specific)	TTGAAGGGCC [A/G] AGGAGATCTG M	A	ט	<u>и</u>
	WIAF-13745	HT27365	3024	PLCB3, phospholipase C, beta 3	GGGCCAAGGA [G/A] ATCTGTTGAA	უ	A	Z Q
	WIAF-13771	HT27365	1079	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	ACATTTTGA [I/C] CCTGAGCAAA S	E	υ	<u> </u>
1	WIAF-13772	HT27365	1546	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	AAGTTGCCTT [C/T] TGATCCAGAT M	υ	E	ν F4
1	WTAF-13773	нт27365	1514	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	MATTABABAG [A/T] ATGATCATTG M	Æ	E	ĸ
i	WIAF-13774	HT27365	1445	PLCB3, phospholipase C, beta 3	AGGICTITGG [C/I] AATAAACICT	υ	E	ŋ
1	WIAF-13778	HT27365	2087	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	TTCATATCAA [G/A]ATCATCAGTG		Ą	×
1	WTAF-13779	HT27365	2367	PLCB3, phospholipase C, beta 3	TGAATGTTTG [C/T] AGCCTGGATA	υ	F	Q
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G2513u14	WIAF-13782	HT27365	Pl 2719	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	CTCATCACCA [G/A] TGACAATACT	U E	Æ	ω .	Z
G2513u15	WIAF-13783	HT27365	P. 2567 (1	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	TTGATGACAT [C/T] TTTAAAATAG	<u>ი</u>	H	н	Н
G2513u16	WIAF-13784	HT27365	P: 2864 (1	PLCB3, phospholipase C, beta 3 2864 (phosphatidylinositol-specific)	TAGAAATGGC [G/A] GACACAGTCC	ა ტ	<b>4</b>	Ą	ď
G2513u17	WIAF-13785	HT27365	2571 (	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	TGACATCTTT [A/T] AAATAGCGGT	N 4	H	×	*
G2513u18	WIAF-13786	HT27365	P 2706 (	PLCB3, phospholipase C, beta 3 (phosphatidylinositol-specific)	TCIGTCATCT [C/T] GGCICATCAC	U E	- E	ద	В
G252u1	WIAF-10195	HT0425	397 B	FCER2, Fc fragment of IgE, low 397 affinity II, receptor for (CD23A)	GAGGGCTGCC [C/T] GGAACGTCTC	<u>Σ</u>	H	ద	×
(425,112	WIAF-10206	HT0425	930 a	FCER2, Fc fragment of IgE, low affinity II, receptor for (CD23A)	ATGGGAGCCA [T/C] GTGGACTACA	S	บ		田
G253u1	WIAF-10175	HT0573	228 £	IFNB1, interferon, beta 1, 228 fibroblast	GGCTTGAATA [C/T] TGCCTCAAGG	ა ა	E	H	Y
G254u1	WIAF-10196	HT0611	466 IL4R,	.L4R, interleukin 4 receptor	TCAGTGCGGA [T/C] AACTATACAC	S	<u>0</u>	А	Д
G254u2	WIAF-10198	HT0611	1474 IL4R,	L4R, interleukin 4 receptor	CATGCCTTCT [T/C] CCACCTTCGG	S	Ü		ᄓ
G254u3	WIAF-10207	HT0611	1902	ILAR, interleukin 4 receptor	AGTGGCTATC [A/G] GGAGTTTGTA	Æ	Ŋ	α	ద
G260u1	WIAF-10186	HT1090	453 t	ILIR1, interleukin 1 receptor, type I	TGTTATAATG [C/G] ACAAGCCATA	υ Σ	<u></u>	4	_D
G261u1	WIAF-10187	HT1101	434	434 IL/R, interleukin 7 receptor	CCTGAGTGTC [A/G] TCTATCGGGA	M	Ü	н	>
G261u2	WIAF-10203	HT1101	517	517 IL7R, interleukin 7 receptor	TTTTAATGCA[T/C]GATGTAGCTT	8	Ü	-=	æ
G267u1	WIAF-11735	HT1877	11.2R 881 beta	IL2RB, interleukin 2 receptor, beta	TCCTCGTGGG [C/T] CTCAGCGGGG	S	H	<u></u> 0	₀

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G267u2	WIAE-II/59	HITG//	010		M	۵ د	, ]
G268u1	WIAF-11758	HT1985	568			E	D
G268u2	WIAF-11734	HT1985	783	783 CD19 antigen	ACGATCGCCC [G/T] GCCAGAGATA	5	4
G270u1	WIAF-11736	HT2415	530	ILGR, interleukin 6 receptor	AGGAGGTGGC [A/G] AGAGGCGTGC	A G A	⋖
G270u2	WIAF-11760	HT2415	1590	1590 ILGR, interleukin 6 receptor	CATTGCCATT [G/A] TTCTGAGGTT M	G A V	н
G270u3	WIAF-11737	HT2415	1510	ILGR, interleukin 6 receptor	CCAGTGCAAG [A/C] TTCTTCTTCA M	D A	Ø
G270u4	WIAF-11761	HT2415	1451	IL6R, interleukin 6 receptor	CTACTAATAA [A/T] GACGATGATA M	A T	z
G270u5	WIAF-11766	HT2415	1843	ILGR, interleukin 6 receptor	TTCCCCAGAT [A/G] GCTGGCTGGG	* Ø	M
G270u6	WIAF-11767	HT2415	1829	1829 ILGR, interleukin 6 receptor	ATACAGACTA [C/T] TTCTTCCCCA	CLL	H
G271u1	WIAF-11762	HT2531	577	CD2, CD2 antigen (p50), sheep red blood cell receptor	TCAGAGGGTC [A/G] TCACACACAA M	T A G I	>
G271u2	WIAF-11739	HT2531	861	CD2, CD2 antigen (p50), sheep red blood cell receptor	GGAAGCCCCA[A/C]CAAATTCCAG M	N N	異
G271u3	WIAF-11768	HT2531	818	CD2, CD2 antigen (p50), sheep red 818 blood cell receptor	CTGGAGACAA [G/A] AGCCCACAGA	Ω α	×
4.1.1.20	WTAR-11738	HT2531	736	CD2, CD2 antigen (p50), sheep red	CCTCTTGATG [G/A] TCTTTGTGGC	۲ . ت . ت	н >
22.7311	WIAF-11763	HT3139	799	IL2RA, interleukin 2 receptor, alpha	ATCATGGTGC [C/T] TGGCTGCCAG	E U	고
G273u2	WIAF-11764	HT3139	956	IL2RA, interleukin 2 receptor, alpha	AAAGTCCAAT [G/C] CAGCCAGTGG	υ υ	Η Σ
2273113	WTAF-11765	HT3139	701	IL2RA, interleukin 2 receptor,	ACGATGACCC [G/A] CCAGAGATCC	S G A	д
7273114	WTAF-11740	HT3139	1133	IL2RA, interleukin 2 receptor, salpha	AAATGACCCA [C/T] GGGAAGACAA	S C	н
4112122	WTAR-11769	HT3139	1163	IL2RA, interleukin 2 receptor,		Q D	
G276u1	WIAF-10192		64,	644 CD4 antigen		ပ ပ	
G276u2	WIAF-10193	HT3670	153	1535 CD4 antigen		. E	ט כ
G276u3	WIAF-10205	HT3670	121	1217 CD4 antigen	TGATGCTGAG [T/C] TTGAAACTGG	ر	1

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227711	WT&F-10007	D10232	851	RENBP, renin-binding protein		CACGTGATTG [A/G] CAAGTTCCTA	M A	ŋ	Д	<b></b>
G277u2	WIAF-10032	D10232	842	RENBP, renin-binding	protein	CTTCGAGCCC [A/G] CGTGATTGAC	Æ	U	ж	_K
G277u3	WIAF-10042	D10232	634	RENBP, renin-binding protein		GCTGGCGGC[A/G]AATACGCAGA	Æ E	ט	м	E
627911	WIAF-10047	K01740	F8 DX 1658 A)	F8C, coagulation factor VIIIc procoagulant component (hemoph A)	r VIIIc, (hemophilia	ACTGATGTCC [G/A] TCCTTTGTAT	<u>ნ</u>	Ø	ద	н
G279112	WIAF-10049	K01740	2328	F8C, coagulation factor VIIIc, procoagulant component (hemophilia A)	_ ~	CCTTACTGAA [G/A] GTTTCTAGTT	დ ე	A	×	×
2279113	WTAF-10050	K01740	4650	FBC, coagulation factor VIIIc, procoagulant component (hemophilia A)		CTGTTCTCCC [G/A] AAACCAGACT	თ დ	A	д	Q.
20 C	WTAF-10061	K01740	6919	F8C, coagulation procoagulant compc A)	r VIIIc, (hemophilia	ccagaagaca [a/g] tgaaagtcac	M	9	Σ	۵
5079115	WIAF-10080	K01740	480	F8C, coagulation factor VIIIc, procoagulant component (hemophi A)	lia	TTAAGAACAT [G/A] GCTTCCCATC	υ Σ	A	Σ	н
9110100	00800 L-R&TW	K01740	2129	FBC, coagulation factor VIIIc, procoagulant component (hemophilia A)		TACATTCTAA [G/A] CATTGGAGCA	<u>ත</u>	4	S	z
CHC 1 25	MTW 10084	K01740	2533	F8C, coagulation factor VIIIc, procoagulant component (hemophilia A)		GTTTGCACAC [A/G] GAACACCTAT	M	හ	ద	_O
G279u8	WIAF-10086	K01740	6639	F8C, coagule procoagulant A)	ution factor VIIIc, component (hemophilia	ACCCICCAAT [I/C]ATIGCICGAI	ω ⊟	υ	н	н
9116752	WIAF-10087	K01740	5957	F8C, coagulation factor VIIIc, procoagulant component (hemophi?)).	ctor VIIIc, nt (hemophilia	GAGAATTATC [G/A] CTTCCATGCA	<u>υ</u>	A	ద	щ
G279a10	WIAF-10495	K01740	F8 pr 5829 A)	F8C, coagulation factor VIIIc procoagulant component (hemophis)A)	ctor VIIIc, nt (hemophilia	TGACAGTACA [G/A] GAATTTGCTC	S D	A	Ø	Q
7279a11	WIAF-10496	K01740	5852	F8C, coagulation factor VIIIc, procoagulant component (hemophi 2 A)	ctor VIIIc, nt (hemophilia	TTTTCACCA[T/G]CTTTGATGAG	Σ	O	н	ß
G279a12	WIAF-10502	K01740	F8 Dr 2492 A)	F8C, coagulation factor VIIIc, procoagulant component (hemoph: 2 A)	ctor VIIIc, int (hemophilia	ACCACAATTC[C/T]AGAAAATGAC	Σ Σ	E-	<u>D4</u>	니

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WIAF-10503 K01740 6906 A   WIAF-13120 K01740 1980 A   WIAF-13121 K01740 1982 A   WIAF-10070 L25615 976   WIAF-10072 L25615 343   WIAF-10072 L25615 343   WIAF-10092 L25615 1075   WIAF-10182 M16827 696   WIAF-10182 M16827 696   WIAF-10182 M16827 696   WIAF-10108 M28372 2558   WIAF-10108 M28372 2558   WIAF-10041 M63012 172   WIAF-10041 M63012 172   WIAF-10041 M63012 172   WIAF-10041 M63012 173   WIAF-10041 M63012 MARTIN M63012 MART	ļ					
WIAF-10503   K01740   6906 A	C, coagulation factor vilit, occoagulant component (hemophilia	אים אימי אים רמי מין אים אים אים איני אים מיות איני אים מיות איני אים מיות איני איני איני איני איני איני איני	ن	E		
WIAF-13120   K01740   1980   A		CAAGTGGA[C/1]11CAGAAGA	_ ر		$\top$	Ţ
WIAF-13120   K01740   1980   A	coagulation factor VIIIc, oagulant component (hemophilia	,				
WIAF-13121   K01740   1982   P   P   P   P   P   P   P   P   P		CAGAGAATAT [A/c] CAACGCTTTC	S	υ	- -1	_
WIAF-13121   K01740   1982   P	coagulation facto		****			
WIAF-13121   K01740   1982 A	occoagulant component (hemophilia	CT: CT: TT: TT: TT: CT: CT: CT: CT: CT:	2	C		Д
MIAF-10067 L25615 976 F MIAF-10070 L25615 460 F MIAF-10071 L25615 343 F MIAF-10072 L25615 535 MIAF-10092 L25615 535 MIAF-10092 L25615 1089 MIAF-10182 MI6827 696 MIAF-1008 MIREZZZZZZSS MIAF-1008 MIAF-10041 MIREZZZZZSS	A)	GAGAATATAC [A/C] ACGCIIICIC		,		
WIAF-10067   L25615   976   E	arginine vasopressin			K	р	-
WIAF-10070   L25615   460   1	1A	CGCCTTTCTT [C/A] ATCATCCAGA	ر ع	∢		,
WIAF-10070   L25615   460   Kar-10071   L25615   343   Kar-10072   L25615   535   Kar-10072   L25615   535   Kar-10092   L25615   L25615	arginine vasopressin	出立しつびつほうつか ログール 単語なる ギンマラケル	E-	ر	[x	[3t.
WIAF-10071 L25615 343 K	1A	100001000 [2 /T] T 10100000	Т		1	
WIAF-10071 L25615 545	arginine vasopressin	GCCIGGCCGA [C/T] CIGGCCGIGG	<u>ი</u>	F	А	Ω
MIAF-10072 125615 68 1	5					
MIAF-10073 L25615 535 1	AVPRIA, arginine vasopiessiii receptor 1A	TCTCTCCGCC [G/A] GTCCCGACGC	Ð	A	Ö	S
WIAF-10073 L25615 535 x wiaf-10092 L25615 1075 x wiaf-10499 L25615 1089 x wiaf-10182 mic827 696 wiaf-10041 m63012 258	K LCCCCCC					
WIAF-10092 L25615 1075 1	receptor 1A	AGACTCTGCA [A/G] CAGCCCGCGC	S S	ŋ	α	a
WIAF-10052 L25615 1089 2 WIAF-10182 M16827 1179 WIAF-10108 W28372 258 WIAF-10041 M63012 172	arginine vasopressin	CCTTGAATAG [C/A] TGCTGTAATC	υ Σ	Æ	S	α.
MIAF-10182 L25615 1089   2						
WIAF-10182 M16827 1179 3 WIAF-10108 M28372 258 WIAF-10041 M63012 172	arginine vasopressin 1A	TGTAATCCCT [G/A]GATATACATG	N D	Æ	×	*
WIAF-10182 M16827 1179 WIAF-10515 M16827 696 WIAF-10108 M28372 258 WIAF-10041 M63012 172	ACADM, acyl-Coenzyme A					
WIAF-10182 MI6827 1179; WIAF-10515 MI6827 696 WIAF-10108 M28372 258 WIAF-10041 M63012 172	C-4 to C-12			ַ	Þ	2
WIAF-10515 M16827 696 WIAF-10108 M28372 258 WIAF-10041 M63012 172		AATATCCTGT [A/G] GAAAAACTAA	V A	5	>	>
WIAF-10515 M16827 696 WIAF-10108 M28372 258 WIAF-10041 M63012 172	acyl-Coenzyme					
WIAF-10515 M16827 696 WIAF-10108 M28372 258 WIAF-10041 M63012 172	C-4 to C-12	C K C C C C FE F C C C C FE C C C C FE C C C C	, c	τ.	<b>4</b>	4
WIAF-10041 M63012 258	straight chain	TTGTGGAAGC [A/ G] GAIACCCCAG		2	¢	\$
WIAF-10108 M28372 258 WIAF-10041 M63012 172	zwro zinc finger protein 9 (a					
WIAF-10108 M28372 258 WIAF-10041 M63012 172	lar retroviral nucleic acid					
WIAF-10041 M63012 172	binding protein)	CTCTTCCAGA [T/C] ATTTGTTATC		<u>U</u>		
LRPAP1, low densit	paraoxonase 1	CTCTGAAGAC [A/T] TGGAGATACT	Ψ W	-	Σ	٦.
LRPAP1, low densit						
	LRPAP1, low density lipoprotein-					
related protein-ass	<pre>slated protein-associated protein (alpha-2-macroglobulin receptor-</pre>					
G290ul WIAF-10085 M63959 354 associated protein	protein 1)	CTCATACGCA [A/G] CCTCAATGTC	Ψ	و	2	n

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			IRPAD1, low density lipoprotein- related protein-associated protein 1 (alpha-2-macroglobulin receptor-	AGCGACTGCA [T/A] CTTCCTCCGG			Q
	WIAF-13122	M63959 M74096	ACADL, acyl-Coenzyme A	AGTGCAACAT [A/C]AATTAGCAGA M	Ω Ω	저	Ø
	00101-1411		LIPA, lipase A, lysosomal acid, cholesterol esterase (Wolman	AAGGACTTAT [T/C] TGGAGACAAA	D E	<u> </u>	ω
	WIAF-10068	M/4//5	LIPA, lipase A, lysosomal acid, cholesterol esterase (Wolman 107 disease)	TGAGGGGTCT [G/A]GAGGGAAACT M	G A	9	<u> </u>
G293a3	WIAF-10498	M74775	LIPA, lipase A, lysosomal acid, cholesterol esterase (Wolman 86 disease)	GGTTCTCTGG[C/A]CCCTGCATTC M	ر ا	Δı	E
G295u1	WIAF-10057	U04270	KCNH2, potassium voltage-gated channel, subfamily H, member 2	AAAGGAGCGA [A/T] CCCACAATGT M	et E	[-1	ഗ
G295u2	WIAF-10062	U04270	KCNH2, potassium voltage-gated 1875 channel, subfamily H, member 2	CGCACTGGGT [A/G]GCCTGCATCT S	4	ر ا	리
G295u3	WIAF-10064	U04270	KCNH2, potassium voltage-gated 2040 channel, subfamily H, member 2	ACTICACCIT [C/I] AGCAGCCICA	υ	[H	[24
G295u4	WIAF-10088	U04270	KCNH2, potassium voltage-gated 1650 channel, subfamily H, member 2	CCGCCCCCAT[C/T]GCCGTCCACT S	U	H	_ <u> </u>
G295u5	WIAF-10090	U04270	KCNH2, potassium voltage-gated 2139 channel, subfamily H, member 2	CCCTCATGTA [T/C] GCTAGCATCT S	H	D N	<b>→</b>
G2951u1	WIAF-14147	HT0030	ZNF42, zinc finger protein 42 (myeloid-specific retinoic acid-1334 responsive)	CCCTGCTCTG [A/G] TCACCACCCG M	٩	<u>н</u>	>
G2951u2	WIAE-14157	HT0030	ZNF42, zinc finger protein 42 (myeloid-specific retinoic acid-1558 responsive)	ACCAGCTTAC [G/A] CACACCGAGG	ტ	4.	<u>E4</u>

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ריים אם כר	WT28-12501	HT0134	1014 I	GRLF1, glucocorticoid receptor	GTGGAGAGAC [T/C] CTGCATAGCT	E	υ	E	E
	7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	нт0134	1853	oid receptor	GAGCCATCTT [A/C] CAGCCTGTTT M	Æ	υ	⊱	S
באפנה	WTAR-10514	U12778	961	me A /branched	TATTCCATAT [A/G] TTAAAGAAAG	Æ	ღ	н	Δ
G2968u1	WIAF-12699	HT0244	1754	SMARCA1, SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 1	CAGAAGAAAC[C/T]AGTACGTGTA M	Ū	타	ф	17
C. S. O. O. C.	777 L	HT0244	2624	SMARCAL, SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily 2624 a, member 1	TGGGAACGTT [G/T] CAATGAATTA M	<u>ن</u>	E	υ	[I4
70000	000000000000000000000000000000000000000	1116660	402	ECH1, enoyl Coenzyme A hydratase	ACATGGCTTC [G/A] GACATCCTGC	<u></u>	Æ	ω	വ
629 /UL	WTAF-10110	016660	149	ECH1, enoyl Coenzyme A hydratase	GCACAAGAGG [A/C] GGCTTCCGGA M	4	υ	回	A
110000	MT&R-12746	HT0281	682	BR140: bromodomain-containing 682 protein, 140kD (peregrin)	atgacatgga [C/T] gaggaggact	<u>ი</u>	E-i	Д	Д
Thou can	WTAE-10709	HT-0334	1104	B-cell-specific transcription	AGITITCCGG [G/A] AGICCCIACA	S G	Æ	U	U
G2975u2	WIAF-12730	HT0334	1185	B-cell-specific transcription 1185 factor	GCTCCCCTA[C/T]TATTAGCG	υ 	E	×	×
G2976a1	WIAF-12129	HT0340	1600	SATB1, special AT-rich sequence binding protein 1 (binds to nuclear matrix/scaffold-associating DNA's)	GTCCTGCCCC [C/A] CTCATCAGCA	ა ა	4	<u> </u>	Сц
G2976u2	WIAF-12743	HT0340	2116	SATB1, special AT-rich sequence binding protein 1 (binds to nuclear matrix/scaffold- 2116 associating DNA's)	TGGCCTCTCC [A/G] GCAGAGTCAG	S A		<u> </u>	<u> </u>
927/045									

G2978ul WIAF-12721 G298ul WIAF-10048 G298u2 WIAF-10051			7 04 L	MSX1, msh (Drosophila) homeo box			E		
		HI 0346	ν Ε Τ Τ	og 1 (formerly homeo box 7)	CATAGAGGGT [C/T] CCAGGTCCCC	اد	H		
G298u2 WIAF-1		U33837	H 36688	Human glycoprotein receptor gp330 8995 precursor, mRNA, complete cds.	CCGGACAGGA [G/A] GTGCATTCCC	<u>ح</u>	4	<u>~</u>	×
		U33837	13217 F	Human glycoprotein receptor gp330 precursor, mRNA, complete cds.	ATGCAGCCAT [C/T] GAACTGCCTA	<u>က</u>	₽	н	н
G298u3 WIAF-10077	10077	U33837	E 6298 I	Human glycoprotein receptor gp330 precursor, mRNA, complete cds.	AACTCTTTCA [T/C] TGTTGTTTCA	E	υ		H
G298u4 WIAF-10078	10078	U33837	6371 [	Human glycoprotein receptor gp330 6371 precursor, mRNA, complete cds.	CCATGGTGCC [G/A] GTGGCAGGCC	ω O	ব	Д	Δι
G298u5 WIAF-	WIAF-10079	U33837	6914 1	Human glycoprotein receptor gp330 6914 precursor, mRNA, complete cds.	ACTCTGAAGT [G/A] ATTCGTTATG	න ව	4	Þ	٥
G298u6 WIAF-10081	10081	U33837	8718	Human glycoprotein receptor gp330 8718 precursor, mRNA, complete cds.	GTTCCAATGC [G/A] CATCTGGGCG	<u>ن</u> ع	A	Æ	<u> </u>
G298u7 WIAF-	WIAF-10083	U33837	9088	Human glycoprotein receptor gp330 9088 precursor, mRNA, complete cds.	ACTTGCTCTG [A/G] AAATGAATTC	Æ	· · · ·	[H	Ü
	WIAF-10096	U33837	6949	Human glycoprotein receptor gp330 grecursor, mRNA, complete cds.	ACTCCTTATG [G/C] CATCACTGTT	<u>υ</u>	U	U	A
	WIAF-10097	U33837	7149	Human glycoprotein receptor gp330 7149 precursor, mRNA, complete cds.	TTGCTTGGAA [A/G] ACAATGGTGG	M	A B	z	
G298u10 WIAF-	WIAF-10100	U33837	8590	Human glycoprotein receptor gp330 precursor, mRNA, complete cds.	TACACAAAAT [G/A] TCATAATTCA	×	<u>ط</u> ق	Ü	
	WIAF-10101	U33837	12948	Human glycoprotein receptor gp330 12948 precursor, mRNA, complete cds.	CAICITIGAA [G/C] ACCAGITAIA	Σ	<u>ပ</u> ဗ		
G2980ul WIAF	WIAF-12723	HT0356	437	TLE1, transducin-like enhancer of split 1, homolog of Drosophila 437 E(spl)	TCATGGCCAC [G/A] GACCCCCAGT	Σ		<u></u>	<u> </u>

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	AGTGGCTGGC [A/G] GTGGGCATGG	CCATGGCAGA [G/A] TTGAATGCCA	ATCGCCAAGA [G/A]ATTGAATACG	GCCACACAGA [C/T] GGAGCCAGCT	CCGCCTGCTA [C/T] GCCCTGGCCA	ACAAATACAT[T/C]GTGACAGGCT	CGGACAGCGT[C/T]GCCCTGAGGA	CTGAGTTGTG [A/T] CATCTCCAGA	TGTCACCCTC [G/C] GAAAGCCTCC	f TGGACAACAC [G/A] GTGCGCTCCT
	TLE1, transducin-like enhancer of split 1, homolog of Drosophila B(spl)	TLE1, transducin-like enhancer of split 1, homolog of Drosophila E(spl)	TLE1, transducin-like enhancer of split 1, homolog of Drosophila (E(spl))	TLE1, transducin-like enhancer of split 1, homolog of Drosophila (6 E(spl))	TLE1, transducin-like enhancer of split 1, homolog of Drosophila 1759 E(spl)	TLE2, transducin-like enhancer of split 2, homolog of Drosophila 06 [E(spl)]	TLE2, transducin-like enhancer of split 2, homolog of Drosophila 36 E(spl)	TLE2, transducin-like enhancer of split 2, homolog of Drosophila [B(spl)]	TLE3, transducin-like enhancer of split 3, homolog of Drosophila 636 [E(spl)]	TLE3, transducin-like enhancer of split 3, homolog of Drosophila 1944 E(spl)
	2044	379	6 276	6 1876		7 2206	1036			
	HT0356	HT0356	HT0356	HT0356	HT0356	) HT0357	HT0357		3 HT0360	4 HT0360
	WIAF-12726	WIAF-12747	WIAF-12748	WIAF-12749	WIAF-12750	WIAF-12720	WTAF-12737	WIAF-12740	WIAF-12833	WIAF-12834
	32980u2	G2980u3	G2980u4	G2980u5	G2980u6	G2981u1	20981u2	G2981u3	G2983u1	G2983u2

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			<u> H                                   </u>	TLE3, transducin-like enhancer of split 3, homolog of Drosophila						
G2983u3	WIAF-12848	HT0360	1710 E		ACCTGGCCTC [G/A] CCCACGCCCC	ν 2	9 0	4 k	מ ב	Т
G2985u1	WIAF-12724	HT0421	995 L	D3	GGCTTCGCCA [G/A] CGCCAACCIG			1 F	7	Т
G2985u2	WIAF-12725	HT0421	1003 h	homeotic protein D3	CAGCGCCAAC [ C/T] TGCAGGCCAG	Ω	ار	<u> </u>	-	Т
1 103860	WIAF-14124	HT0468	1197 CSDA,	cold shock domain protein A	GCCGTGGATA [C/T] CGGCGTCCCT	ß	υ	T	7	
	WTAF-12758	HT0474	2068 4	ZNF7, zinc finger protein 7 (KOX 4, clone HF.16)	AGTGGTTTTA [C/T] GAATATGGGA	S	U	T	- 7	
	WTAF-12773	HT0474	985 4	ger protein 7 (KOX	GAGAGAAGCC [G/C] TACGAATGTG	တ	Ü	С	Д	
	TOUCH SET ME	HT0474	1278	ger protein 7 (KOX	AGCCAGCAGT [C/T] GCAGCTGGTT	Σ	υ	ω Ε		
	WIAF-12133	HT0735	1441	n 5.1	GAGGCAGCGG [C/T] CCCGGGCCTG	S	υ	EH	D D	Т
G3008a1	WIAF-12134	HT0753	1850	ATF4, activating transcription factor 4 (tax-responsive enhancer 1850 element B67)	Taaaagag [g/a] gcggaittccc	w	ט	4	~ ~	
G3008u2	WIAF-12798	HT0753	946	ATF4, activating transcription factor 4 (tax-responsive enhancer element B67)	CCCTTCGACC[C/A] GTCGGGTTTG	Σ	υ	Æ	д	õ
3300813	WIAF-12812	HT0753	1482	ATF4, activating transcription factor 4 (tax-responsive enhancer 1482 element B67)	CACTGCTTAC [G/A] TTGCCATGAT	Σ	Ð	A	<b>D</b>	Н
G3008u4	WIAF-12813	HT0753	1847	ATF4, activating transcription factor 4 (tax-responsive enhancer element B67)	CTCTAAAAGA [G/C]AGGGCGGAIT	Σ	ט	Ü	[2]	Д
G301u1	WIAF-10127	U71285	3639	MTR, 5-methyltetrahydrofolate- homocysteine methyltransferase	TGTGGAGACT [C/T] GCAGACATCG	တ	υ	H	리	LI .
G3012u1	WIAF-12794	HT0873	402	402 MAX dimerization protein	TGGTGCCACT [G/T] GGACCCGAAT	ω	υ	H	н	ı,
G3014u1	WIAF-14183	HT0899	274	274 homeotic protein 2, distal-less	AAAAGACTCA[G/A]TACTTGGCCT	ω	_O	Æ	Ø	a

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	<u></u>			MLLT3, myeloid/lymphoid or mixed- lineage leukemia (trithorax			
G3.02.0m1	WIAF-12797	HT0956	852	(Drosophila) homolog); translocated to, 3	GIGCCITCAA [A/G] GAACCITCCA	S A G	×
5 5	WT&R-13724	HT0966	381	zinc finger, X-linked, duplicated A	GCTGCAGCAA [G/A] CAATATGACA	S G A K	×
63023UI	FO/OT TUTM			zinc finger, X-linked, duplicated			
G3023u2	WIAF-13725	HT0966	220 A		GGCCAAACTC [G/A] GCGCCCACCA		1
23023113	WIAF-13726	HT0966	69	zinc finger, X-linked, duplicated A	AGTCGCACGA [T/C] AAACTGCGGC	STCD	Д
	MINE-12727	нточк	Z49 A	zinc finger, X-linked, duplicated	ACTTCGAACC [C/T] GAGAGGCCTT	S C	<u>д</u>
65025u <del>1</del>	TOTAL TOTAL			zinc finger, X-linked, duplicated	CABECAGTAG	M G A	A T
G3023u5	WIAF-13765	HT0966	T99				
31152025	WTAF-13766	HT0966	1302 A	ınc ıınger, A-ıımveu,	TGACTCCTTC [G/T] AGCACCCTTT	E U	
G302343	WTAF-12800	HT1035	124	124 HOXB7, homeo box B7	TTATGCGAAT [G/A] CTTTATTTTC	G	- 1
23027112	WTAF-12816	HT1035	450	450 HOXB7, homeo box B7	GGGACTCGGA [C/T] TTGGCGGCCG	L L	
G3028u1	WIAF-12806	HT1037	701	701 homeotic protein C8	AGACCCTGGA [A/G] CTGGAGAAGG	ن ا	
200000	WT&F-14153	HT1100	441	441 zinc finger protein 8	TCAGACTCAG [G/A] GAAAACTGCG	G	
G3029u2	WIAF-14155	HT1100	1416	1416 zinc finger protein 8	GGCGTGAACA [A/G] TCCTCGAGCA	S A G	a a
	C C C C C C C C C C C C C C C C C C C	71001	0.110	LRP1, low density lipoprotein- related protein 1 (alpha-2-	ATGGAGCTGG [G/A] GCCCGACAAC	M G A	<u>ස</u>
G303u1	WIAF-LUUUU	PTCCTV	7	1			
G303u2	WIAF-10001	X13916	4012	LRP1, low density lipoprotein- related protein 1 (alpha-2- 4012 macroglobulin receptor)	GCGAGCTCTG[C/T]GACCAGTGCT	S S	U U
G303n3	WIAF-10002	X13916	4702	<pre>LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)</pre>	GCCIGCCCG [C/T] ATTGAGGCAG	H C N	ద
G303u4	WIAF-10003	X13916	6395	LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)	CTGGATCGCA [G/A] GCAACATCTA	M G A	ධ ග
203115	WIAE-10004	X13916	6937	LRP1, low density lipoprotein- related protein 1 (alpha-2- 6937 macroglobulin receptor)	AAGGCACCAA [C/T] GTGTGCGCGG	S C D	Z Z

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G303u6 W	WIAF-10005	X13916	I x 9391	LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)	GGCTGAAGGA [T/C] GACGGCCGGA	E	ט	<u> </u>
	WIAF-10011	X13916	766 1	LRP1, low density lipoprotein- related protein 1 (alpha-2- 766 macroglobulin receptor)	ACTGCATGGA[C/T]GGCTCAGATG	υ	F	О
6303u8	WIAF-10015	X13916	9040	LRP1, low density lipoprotein- related protein 1 (alpha-2- 9040 macroglobulin receptor)	ACCCGACCTG [C/T] GGCCCCAGTG S	U	H	υ υ
630309	WIAF-10019	X13916	11749	LRP1, low density lipoprotein- related protein 1 (alpha-2- 11749 macroglobulin receptor)	CCCTGCGCTG [C/T] AACATGTTCG	ე დ	E	ט
G303u10	WIAF-10020	X13916	1917	LRP1, low density lipoprotein- related protein 1 (alpha-2- 1917 macroglobulin receptor)	GACCAGTATG [G/A] GAAGCCGGGT	<u>ა</u>	Æ	<u>м</u>
G303u11	WIRE-10021	X13916	4810	LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)	AGAAGCGCAT [C/T] CTTTCGATTG	υ ω	E	H
	WIAF-10022	X13916	6367	LRP1, low density lipoprotein- related protein 1 (alpha-2- 6367 macroglobulin receptor)	TTGGCCGTGT [G/C] GAGGGCAFTG	<u>හ</u>	ט	Δ Δ
G303u13	WIAF-10023	X13916	6247	LRP1, low density lipoprotein- related protein 1 (alpha-2- 6247 macroglobulin receptor)	CTGTCGGCAT[C/T]GACTTCCACG	<u>ე</u>	H	н
G303u14	WIAF-10024	X13916	8371	LRP1, low density lipoprotein- related protein 1 (alpha-2- 8371 macroglobulin receptor)	acgcctcaga [t/c] gagatgaact	رم ا	U	<u>а</u>
G303u15	WIAF-10030	X13916	11395	LRP1, low density lipoprotein- related protein 1 (alpha-2- 11395 macroglobulin receptor)	ACGGCAGCGA [C/T] GAGGAGGCCT	<u>ა</u>	E	Д

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G303u16	WIAF-10031	X13916	12763 11	LRP1, low density lipoprotein- related protein 1 (alpha-2- 12763 macroglobulin receptor)	ACGICITIGA [G/A] GALIACAICT	d G	ы ы	
G303u17	WIAF-10035	X13916	640 n	LRP1, low density lipoprotein- related protein 1 (alpha-2- 640 macroglobulin receptor)	ACGGATCTGA [C/T] GAGGCCCCTG	Ð	D D	
G303u18	WIAF-10037	X13916	16091	LRP1, low density lipoprotein- related protein 1 (alpha-2- 1609 macroglobulin receptor)	GCCGCCTTGT [C/T] TACTGGGCAG	EI U	> >	
G303u19	WIAF-10038	X13916	1629	LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)	GATGCCTATC [T/G] GGACTATATT M	υ Ε-	Li Li	2
G303u20	WIAF-10039	X13916	2210	LRP1, low density lipoprotein- related protein 1 (alpha-2- macroglobulin receptor)	CACCAGCTAC [C/T] TCATTGGCCG	E U		E4
G303u21	WIAF-10043	X13916	7287	LRP1, low density lipoprotein- related protein 1 (alpha-2- 7287 macroglobulin receptor)	GATGGCTCCA [G/A] GAGGATCACC	ط ق ع	ᄶ	×
G303u22	WIAF-10044	X13916	8258	LRP1, low density lipoprotein- related protein 1 (alpha-2- 8258 macroglobulin receptor)	CTCTGACGAG [A/G] TCCCTTGCAA	В В	Н	Λ
G303u23	WIAF-10045	X13916	11871	LRP1, low density lipoprotein- related protein 1 (alpha-2- 11871 macroglobulin receptor)	GTGCGCACCG [A/G] GAAAGCGGCC	ტ 4	М	_U
G3031u1	WIAF-14097	HT1128	611	PSMC3, proteasome (prosome, macropain) 26S subunit, ATPase, 3	TGGGGATCCA [A/G] CCTCCAAAG	ν 4	Q	a
G3034u1	WIAF-12836	HT1182	137	TCF12, transcription factor 12 (HTF4, helix-loop-helix 137 transcription factors 4)	ATAAGGGAGC [G/A] TGAGGAGTCT	M G	ਲ	н

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ATCTTCAATT [A/G] TGGGTTCCTT	agagaaggct [a/g] tgcagcttgc	TGTACCAGAC [G/A] CCCTTGCACT	AGCTGCAGCT [G/C] TATAAGTTAC	AACAGCCCCG [G/T] AAGTGGCACC	CGCCAAATGA [G/T] TCAGCTGGCA	טטקטקטקע נט/ צן ע עוויטטטעטטטר	GTGCCGAGGG [G/A] CGGCCACACT	TCCGTTTCCT [C/T] GAGAGCCTGC	GGATTAAGAA [G/A] GCAGCCGAAG	TCCAAGAAGA [T/C] GAAATTCCAG	AGIGGAGCGI [C/ 1] GCCGCCGAGA	CCCTTGTCAT [C/T] GAGTTCACG
TCF12, transcription factor 12 (HTF4, helix-loop-helix transcription factors 4)	NFKB1, nuclear factor of kappa light polypeptide gene enhancer in 1700 B-cells 1 (p105)	NFKB1, nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (p105)	NFKB1, nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (p105)	<pre>GLI3, GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly syndrome)</pre>	<pre>GLI3, GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly syndrome)</pre>	FABP3, fatty acid binding protein 3, muscle and heart (mammary-	158 derived growth inhibitor; IRF2, interferon regulatory 842 fartor 2		1746 transcription factor 1, nucleolar	1829 transcription factor 1, nucleolar	628 transcription factor USF	prolyl 4-hydroxylase, beta subunit/protein disulfide isomerase/thyroid hormone-binding
421 t	1700	1936	2641	3761	3963		158	1233	1746	1829	628	777
HT1182	HT1373	HT1373	HT1373	HT1375	HT1375		HT637	HT1518	HT1518	HT1518	HT1530	HT0034
WIAF-12837	WIAF-12864	WIAF-12881	WIAF-12882	WIAF-13027	WIAF-13028		WIAF-12242	WIAF-1280/ WIAF-12875	WIAF-12876	WIAF-12877	WIAF-12884	WIRF-10150
G3034u2	G3038u1	G3038u2	63038113	G3039n1	63039112		G304u1	G3043u1 G3047u1	G3 04 7u2	G3047u3	G3048ul	G305u1

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G305u2	WIAF-10154	HT0034	1861	prolyl 4-hydroxylase, beta subunit/protein disulfide isomerase/thyroid hormone-binding 186 protein, alt. transcript 1	TGGCGGCCCA[C/A]AAGTACCTGC M C	4	н н	ο
G305u3	WIAF-10155	HT0034	1428	prolyl 4-hydroxylase, beta subunit/protein disulfide isomerase/thyroid hormone-binding protein, alt. transcript 1	GGACGGTCAT [T/C] GATTACAACG S T	ט	н	н
G3050u1	WIAF-12860	HT1558	2098	FSRG1: female sterile homeotic- 2098 related gene 1 (mouse homolog)	AACATTGCAA[T/C]GGCATTTTGA S T	ט	z	
G3050u2	WIAF-12861	HT1558	2845	FSRG1: female sterile homeotic- related gene 1 (mouse homolog)	TAGGCCCTTC[T/C]GGCTTTGGAC S T	<u>U</u>	Ŋ	Ø
G3050u3	WIAF-12862	HT1558	3409	FSRG1: female sterile homeotic-3409 related gene 1 (mouse homolog)	CCTCGTCGTC[G/A]TCTTCAGACA S G	A	w	w_
G3050u4	WIAF-12874	HT1558	1699	FSRG1: female sterile homeotic- 1699 related gene 1 (mouse homolog)	TCTCTTCTGT [G/C] TCACACACAG S G	<u>U</u>	<b>&gt;</b>	>
G3050u5	WIAF-12878	HT1558	2093	FSRG1: female sterile homeotic-related gene 1 (mouse homolog)	GTTAAAACAT[T/G]GCAATGGCAT M T	<u>છ</u>	Ü	ro l
G3050u6	WIAF-12879	HT1558	2746	FSRG1: female sterile homeotic- 2746 related gene 1 (mouse homolog)	CTGGGGCCGA [C/T] GAAGATGACA	U E	Д	
23051111	WTAF-12866	HT1569	1423	MEF2B, MADS box transcription enhancer factor 2, polypeptide B (myocyte enhancer factor 2B)	CITGGCCGAC [G/A] GCTGGCCCCG	ط ق	E-1	H
2.1.1.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2	WT&F-13022	H71369	661	MEF2B, MADS box transcription enhancer factor 2, polypeptide B (myocyte enhancer factor 2B)	CAGAGTACAG[C/T]GAGCCCCACG	H ت	<u> </u>	ω
G2057a1	WIAF-12142	HT1669	5565		AGACTGCTCT [T/C] GAGGCTCATA S	EH	U L	ᆈ
G3057a1	WIAF-12143	HT1669	5634	alpha-fetoprotein enhancer-binding 5634 protein	CTCTGTCTGC [G/A] ATGCTCTTAG S	ט	4	
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			10	alpha-fetoprotein enhancer-binding	GGGGACTCCA [G/T]ATGAAAGGAG	<u>5</u>	E→	0	н	<del></del>
G3057a3	WIAF-12144 WIAF-12145	мт1669		etoprotein enhancer-binding	GCTTTTCCCA [C/T] CTACCCCCAA	υ υ	<u>-</u>	#	異	<del></del> 1
230200	WTAF-12885	HT1669		etoprotein enhancer-binding	TCTGGAGATC [C/T] ATATGAGGTC	Σ U	E-1	田	_ <del>&gt;</del> -	- 1
3172025	WIAF-12892	HT1669		alpha-fetoprotein enhancer-binding	AGACCTTGCC [G/A] GCTCAGCTAC	S	Æ		Δı	<del>-  </del>
23.05.21.7	WTAF-12893	HT1669	4137	etoprotein enhancer-binding	CAAGGTTTAC [G/A] GACTACCAGC	ω υ	Æ	H	H	<del></del>
23057118	WIAF-12897	HT1669	4783	alpha-fetoprotein enhancer-binding protein	GAAGACCAAC [A/C] CTCCCCAGCA	Æ	U	H	д	
230520	WIAF-12898	HT1669	5215	alpha-fetoprotein enhancer-binding 5215 protein	TCCAACCTCC [A/C] CAATGAACAC	M	U	E	Д	
G3057u10	WIAF-12904	HT1669	7266	alpha-fetoprotein enhancer-binding	CCCTGCAGGC [C/T] GCGTTGACTT	ω Ω	E-	A	Æ	
3057111	WIAF-12907	HT1669	8345	alpha-fetoprotein enhancer-binding 8345 protein	CCAACAGACG [A/C] CTATTCGGAG	M	0	Ω	< 4	
G3057u12	WIAF-12943	HT1669	4257	alpha-fetoprotein enhancer-binding protein	TGGTGTGGTT[T/C]CAGAATGCCC	S	υ E.	Γtι	ഥ	1
2000	MTAH-12951	HT1669	7333	alpha-fetoprotein enhancer-binding protein	ACCAGGCTTT [T/A] CTCCTTATTA	Σ	A	Ø	E-I	<del></del>
45.000 41.1740 41.1740	WTAF-13030	HT1669	303	alpha-fetoprotein enhancer-binding protein	GCAGCCTGTC [G/A] GAGGACGAGT	တ	<b>₹</b>	S	တ	
4 1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	WTAR-13031	HT1669	777		GCCTTCCAGA [G/A] GAGGACGAGG	တ	ري ن	E4	[23]	
1000	α	нт0040	1618	CPT2, carnitine	CTCTACTGCC [G/A] TCCACTTTGA					
G307111	WIAF-10076	HT0114	110	110 EDN2, endothelin 2	CGTTGCGCTA [G/A] CCCTGCTCGT	Σ	8	A	⊢	$\neg$
	WIAF-12972	HT2085	625	pre-B-cell leukemia transcription factor 3	agaaatatga [a/g] caggcatgta	ß	4	D E	E	
G3070u2	WIAF-12973	HT2085	841	pre-B-cell leukemia transcription factor 3	GTAACTTCAG [T/C] AAACAGGCCA	ω	ы	ري د	S	
G3071u1	WIAF-12886	HT2086	995	AGER, advanced glycosylation end product-specific receptor	CCTGCGAGGC[T/C]GTGATGATCC	Ø	H		4	
G3071u2	WIAF-12887	HT2086	1475	AGER, advanced glycosylation end product-specific receptor	GAGGCCAGAT [C/G] TACAGCCCAC	Σ	U	 	Σ	
G3071u3	WIAF-12935	HT2086	933	AGER, advanced glycosylation end 933 product-specific receptor	ACGCATGGTG [A/G] GCATCATCCA	Σ	A		<u>ი</u>	

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G3071u4	WIAF-12936	HT2086	1052	AGER, advanced glycosylation end 1052 product-specific receptor	GTAACTTCAG [C/T] AAACAGGCCA	S C	E	ω	co
31115	WT 2 H - 1 2 9 3 7	HT2086	8361	AGER, advanced glycosylation end 836 product-specific receptor	agaagtatga [g/a] caggcatgta	<u>ი</u>	4	[E]	凶
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G308U1	WIAF-1005*	20 C C C C C C C C C C C C C C C C C C C	33 88		GGGATGATGA [C/T] GCCCACGGTG	υ Σ	E	Ħ	Σ
65000L2	WTAR-12997	HT2188	689	PSMC2, proteasome (prosome, 689 macropain) 26S subunit, ATPase, 2	GCCATTGAGC [C/T] TCCCAAGGGC	<u>ت</u>	E		Ы
13.400.00	WTAF-12976	HT2228	106	IGHMBP2, immunoglobulin mu 106 binding protein 2	TGCTGGAGCT [T/C] GAGAGAGACG	S	υ	ᄓ	'n
2300000	WIAF-12985	HT2228	2260	IGHMBP2, immunoglobulin mu 2260 binding protein 2	TGGAGTTCAT [G/C] GCCAGCAAGA		U	Σ.	н
23083113	WIAF-12986	HT2228	2060	IGHMBP2, immunoglobulin mu 2060 binding protein 2	GGGACCTGCT [A/G] CGTCCACCAG	M	Ü	٤٠	Æ
7:0000000000000000000000000000000000000	WTAR-12987	HT2228	2365	IGHMBP2, immunoglobulin mu 2365 binding protein 2	ACGACAGTTC[C/T]GGGGAAGGGA	S C	H	တ	ß
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9468065	MTAF-13006	HT2228	272	IGHMBP2, immunoglobulin mu 272 binding protein 2	ATACGGGTCC [G/A] CGGCAGCTCT	ַ ב	4	Æ	E
G3083117	WIAF-13010	HT2228	2581	IGHMBP2, immunoglobulin mu 2581 binding protein 2	TCAGGAGCGC [G/A] CAGGGGCAGC	S D	4	Æ	Æ
G3083u8	WIAF-13011	HT2228	2594	IGHMBP2, immunoglobulin mu 2594 binding protein 2	GGGGCAGCCC [G/A] CCAGCAAGGA	Σ U	A.	₹.	H
G3088u1	WIAF-12984	HT2318	884	HIVEP1, human immunodeficiency virus type I enhancer-binding protein 1	TGTGGCACTA [C/T] GTCCCCCTCC	Σ Σ	E+	EH	Σ
G3088u2	WIAF-12988	HT2318	2469	HIVEP1, human immunodeficiency virus type I enhancer-binding 2469 protein 1	TCTTGTCACC[A/G]CGTCAACACC	8	A G		Δı

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HIVEFI, numan immunodeliciency virus type I enhancer-binding 3066 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 4008 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding	HIVBP1, human immunodeficiency virus type I enhancer-binding 5148 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 5834 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 6065 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 7652 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 741 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 948 protein 1	Vinetinia Comming to the transmission of the contract of the c
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TCTTCTGTCT [G/A] TACCTTCACT	GCGGTCTGCA [A/G] CCTCAGATTC	CCTAAACATA [G/A] TGTTACCATA	TGGGTCTTCT [A/G] AAAGTGAGGA	CCGCTCTGTA[C/T]ACCTGGTACG	eggccgagcc[c/T]gacaccaagc	CCAGTTCTCC [C/T] AGCAGCTGCA	GAATGAGTGT [A/G] AGTTGCAGAA	CGCCTTCTTC [G/T] CCCGAGGACA	CCTTGGACCA [G/T] CTCATCCAGA	CTGAGGACAA [G/A] GCCAACAAGA	GTCCCTGTTA [C/T] GGCTACCAGT	TCATATTCAT [C/T] AGAGGAAATG	TACTCCAGAG [G/A] TCAAGTCCAA
HIVEP1, human immunodeficiency virus type I enhancer-binding 2803 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 3342 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 542 protein 1	HIVEP1, human immunodeficiency virus type I enhancer-binding 4972 protein 1	TCF2, transcription factor 2, hepatic; LF-B3; variant hepatic 701 nuclear factor	TCF2, transcription factor 2, hepatic; LF-B3; variant hepatic 362 nuclear factor	TCF2, transcription factor 2, hepatic, LF-B3; variant hepatic	ZNF141, zinc finger protein 141 526 (clone pHZ-44)	NRF1, nuclear respiratory factor 259 1	1106 E2F2, E2F transcription factor 2	1154 E2F2, E2F transcription factor 2	1339 A2M, alpha-2-macroglobulin	1201 A2M, alpha-2-macroglobulin	3041 A2M, alpha-2-macroglobulin
2803	3342	3542	497	702	36	69	52	25	110	115	133		30,
HT2318	HT2318	HT2318	HT2318	HT2435	HT2435	3 C L C THE	HT2483	HT2508	HT2511	HT2511	HT0402	HT0402	HT0402
WIAF-13013	WIAF-13015	WIAF-13016	WIAF-13017	WIRE-12994	WIAF-13018		WIAF-13020 WIAF-12147	WIAF-12975	WIAF-13617	WIAF-13659	WIAF-10291	WIAF-10292	WIAF-10293
23.0821.13	G3088u14	G3088u15	G3088u16	G3095u1	63095112		G3095u3	G3102u1	G3103u1	G3103u2	G311u1	G311u2	G311u3

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G311u4	WIAF-10294	HT0402	3676 A2M,		alpha-2-macroglobulin	TGACATCCTA [T/C] GTGCTCCTCG	တ	E	ט	7	×
G311u5	WIAF-10296	HT0402	3364 A2M,		alpha-2-macroglobulin	ATATCACCAT[C/T]GCCTTCTGG	w	υ	E	Н	н
G311u6	WIAF-10297	HT0402	3203 A2M,		alpha-2-macroglobulin	CCAAGCTCGA [G/T] CCTACATCTT	×	_o	E	4	ω
G311a7	WIAF-10494	HT0402	1122 A2M,		alpha-2-macroglobulin	TCACACTTTC[G/A]ACAGGAATT	Σ	_O	A	24	a
G3119u1	WIAF-13947	HT2654	2876	GLI, glioma	GLI, glioma-associated oncogene 2876 homolog (zinc finger protein)	TTTCIGGGG [G/A] TTCCCAGGTT	Σ	U	A	ש	Ω
G3119u2	WIAF-13959	HT2654	654 1	GLI, gliom homolog (zi	GLI, glioma-associated oncogene 654 homolog (zinc finger protein)	AGTGCCGGGA [G/A]GAACCCTTGG	w	U	Æ	E	闰
23119113	WIRF-13965	HT2654	3376 1	GLI, gliom homolog (zi	GLI, glioma-associated oncogene 3376 homolog (zinc finger protein)	TGGGGAAACA [G/C] AAITCCICAA	Σ	ŋ	Ü	ы	ø
G31211	WIAE-10006	HT0428	86.8	PLAU, plas	plasminogen activator, se	CTCACCACAA [C/T] GACATTGCCT	ß	υ	E-1	Z	z
G312u2	WIAF-10029	HT0428	498	PLAU, plas 498 urokinase	plasminogen activator, ase	GGCCTAAAGC [C/T] GCTTGTCCAA	Σ	ŭ	Е	Дı	ı
G312a3	WIAF-10521	HT0428	767	PLAU, plas	plasminogen activator, use	TGATTACCCA [A/C]AGAAGGAGGA	Σ	Æ	υ	M	a
G3125u1	WIAF-13675	HT2674	740	GTF2F2, ge factor IIF, 740 subunit)	GTF2F2, general transcription factor IIF, polypeptide 2 (30kD subunit)	acatcacaaa [a/g] caacctgtgg	S	æ	v	×	×
G313u1	WIAF-10129	HT0462	3086	platelet-de alpha poly	platelet-derived growth factor, 3086 alpha polypeptide (GB:M21574)	CATGCGTG[G/A]ACTCAGACAA	Σ	Ö	Æ	А	z
G313u2	WIAF-10130	HT0462	1078	platelet-de alpha poly	platelet-derived growth factor, 1078 alpha polypeptide (GB:M21574)	ATGAGAAAGG [T/G] TTCATTGAAA	ß	H	_O		υ
G313u3	WIAF-10133	HT0462	1571	platelet-derived g alpha polypeptide	platelet-derived growth factor, alpha polypeptide (GB:M21574)	GGAGATCCAC [T/C] CCCGAGACAG	Σ	E	U	လ	Δ,
631304	WIAF-10135	HT0462	2611	platelet-derived g alpha polypeptide	platelet-derived growth factor, alpha polypeptide (GB:M21574)	CTCGCAACGT [C/T] CTCCTGGCAC	S	υ	E+	>	>
G314u1	WIAF-10069	HT0467	1890	ALOX15, arad 1890 lipoxygenase	arachidonate 15- ase	TCAGGGAGGA [G/A] CTGGCTGCCC	ω	ರ	⊿	斑	ഥ

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23141111	WTAF-13934	HT27498	878	NFATC3, nuclear factor of activated T-cells, cytoplasmic 3	CCAGAGGATA[G/A]CTGGCTACTC M G	4	Ø	Z
0014100 0117100	WTAF-13936	HT27498	1189	NFATC3, nuclear factor of activated T-cells, cytoplasmic 3	GCCTGCCTCA[T/C]GCAATGGGAA M T	υ -	υ	P.
G3141u3	WIAF-13938	HT27498	2241	NFAIC3, nuclear factor of 2241 activated T-cells, cytoplasmic 3	CTCTGCGGGG [T/C] TTCCCTTCAG S T	<u></u>		ರ
G3141n4	WIAF-13944	HT27498	702	NFATC3, nuclear factor of activated T-cells, cytoplasmic 3	ATGCCTCTGA[C/T]GAGGCAGCCC S C	H	<u> </u>	
G3159u1	WIAF-13891	HT2757	523	SP4, Sp4 transcription factor	CTTCAAAAGA[G/A]AATAACGTTT S G	A .	EX	<u> </u>
G3159u2	WIAF-13892	HT2757	1514	SP4, Sp4 transcription factor	ACAGAATGTT [C/T] AACTTCAAGC N C	E1	O/	*
23159113	WIAF-13893	HT2757	2236 SP4,	SP4, Sp4 transcription factor	တ			- 1
G3165u1	WIAF-13860	HT27636	437	437 transcription factor B-ATF	AGCAGCTCAC [A/G] GAGGAACTGA S A	ט ט	Δ μ	- Δ
G3165u2	WIAF-13861	HT27636	512	ription factor B-ATF				
G3173u1	WIAF-13556	HT2772	1686	ZNF74, zinc finger protein 74 (Cos52)	TGCACAGCGA [G/A] GGGAAGCCCT S G	<u>ا</u> که	[2]	田
23175111	WTAR-13948	HT2776	2037	transcriptional regulator, via 2037 glucocorticoid receptor	TGTTCGGACC [A/G] GAAGCACCCA S A		D D	Дı
1,000	WT2F-14036	HT2783	1614	MHC2TA, MHC class II 1614 transactivator	ATCCTAGACG[C/G]CTTCGAGGAG M C		رط 20	ט
G3182112	WIAF-14037	HT2783	2791	MHC2TA, MHC class II transactivator	TGAGCGACAC [G/A] GTGGCGCTGT S G		A	E4
2318213	WTAF-14059	HT2783	1657	MHC2TA, MHC class II 1657 transactivator	TGCACAC[G/A] TGCGGACCGG S G	Ü	T A	[-
23182114	WIAF-14060	HT2783	1606	MHC2TA, MHC class II transactivator	TTCTGCTCAT[C/T]CTAGACGCCT S C	Ü	H	н
G3183u1	WIAF-13950	HT27861	392	zinc finger protein C2H2-150	TACTCTAGAG [G/A] AGCCTGTTGG M G	_O	A H	- ×
G3184u1	WIAF-13864	HT27862	271	271 zinc finger protein C2H2-171	GAAACTCCAG [T/G] TCAAAGACTT M T	E	ro O	FI >
G3184u2	WIAF-13865	HT27862	248	zinc finger protein C2H2-171	CIGCTIGAAT [1/C] CAIGTAIGAR M I	ы	ט	Ľτ _ι Ο
G320u1	WIAF-10136	HT0791	552	552 ANX7, annexin VII (synexin)	CCAACTTCGA[T/C]GCTATAAGAG S T	E	υ υ	D

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	70101 HWTM	HP0791	1350 ANX7. annexin VII (synexin)	TTGACCTTGT [A/G] CAAATAAAAC	A G	Þ	Δ
G32002	WIAF-14186	HT27930	£i	GTCAGAAGTC [A/G] GCCCTAATTG S	A G	S	ഗ
G3218u1	WIAF-13526	HT28104	zinc finger protein ZNF169, 187 Krueppel-type	CCCGACAGCT [C/T] ATTAAGAAAG M	٦ ابا	н	¥
	33001-541W	HT0015	Homo sapiens inducible nitric oxide synthase (NOS) mRNA, 1361 complete cds.	ACTICIGIGA [C/I] GICCAGCGCT S	D D	Ω	Д
6323ur	OCCUPATION OCCUPATION	2700HH	FBN1, fibrillin 1 (Marfan 3817 syndrome)	TGTGAATGCC[C/T]GCCTGGCCAT M	<del>ا</del> ن	д	Ы
G325u1	00101-3414		FBM1, fibrillin 1 (Marfan	AGATAGCTCC [T/G] TCCTGTGGCT	<u>ნ</u>	Δ4	д
G325u2	WIAF-10113	79601H		GATCTGCAAT [A/C] ATGGACGCTG M	٦ 2		H
2222002	MT2E-10116	2960шн	FBN1, fibrillin 1 (Marfan 3603 syndrome)	GAACTGCACA [G/C]ACATTGACGA M	ນ ຫ		田
G325u <del>s</del> G325u5	WIAF-10117	HT0962		TCTGCATGAA[C/T]GGGCGTTGCG	EH U	z	Z
G326u1	WIAF-10036	HT1009	KLKB1, kallikrein B plasma, 1854 (Fletcher factor) 1	GCAAACACAA [C/T] GGAATGTGGC	υ L	z	z
G327u1	WIAF-10052	HT1011	1599 HRG, histidine-rich glycoprotein	AAGCCAGACA [A/T] TCAGCCCTTT M	A		н
012000	WTAF-10054	HT1011	1083 HRG, histidine-rich glycoprotein	CCACTATTGC [C/T] CATGTCCTGC M	E U	Дı	니
6327 uz	7 2 0 0 1 - H & T W	HT1011	1140 HRG, histidine-rich glycoprotein	GCCCAAAGAC [A/G] TTCTCATAAT			잱
G328u1	WIAF-10145	HT1087	255 SAA1, serum amyloid A1	GTGCCTGGGC [T/C] GCAGAAGTGA		T	4 :
G328a2	WIAF-10511	HT1087	248 SAA1, serum amyloid A1		<b>-</b>	<b>x</b> :	2 ح
G328a3	WIAF-10512	HT1087	305 SAA1, serum amyloid A1	TTCTTTGGCC [A/G] TGGTGCGGAG M	5 C 4 E		4 1
G328a4	WIAF-13126	HT1087	SAA1, serum amyloid				1 0.
G328a5	WIAF-13127	HT1087	82 SAA1, serum amyloid A1	CINGGICCIG [6/A] GIGICAGCAG	T		2
G329u1	WIAF-10140	HT1141	PLCG1, phospholipase C, gamma 1 2514 (formerly subtype 148)	CIGACCTICA[I/C]CAAGAGCGCC M	L L	H	E
G329u2	WIAF-10162	HT1141	PLCG1, phospholipase C, gamma 1 1036 (formerly subtype 148)	TATGCCCGGA [C/A] ACCATGAACA M	۸ 2		M
G329u3	WIAF-10163	HT1141	PLCG1, phospholipase C, gamma 1 911 (formerly subtype 148)	GITCAIGCIC [A/G] GCITCCICCG	A G	υ O	<u></u>

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G3295u1	WIAF-14017	HT3460	1229 E	FUBP, far upstream element 1229 binding protein	CCATAAAAG [C/T] ATAAGCCAGC	ა ე	E	တ	ß
G3296u1	WIAF-14168	HT3466	6289 E	transcription factor TFIIIC, RNA 6289 polymerase III, alpha subunit	CAGCCTGGAC [G/A] AGAGCCCCAT	υ Σ	A	[2]	×
( (	0.000	и 112766	23.5 T	transcription factor TFIIIC, RNA	GGGCATCAGC [T/A] TCTATGAGGA	E H	4	ļīz.	н
G3296u2	WIAF-141/9	111.3±00	1803		ACTTTGCCAA [C/T] GTGCAGGAGC	S C	H	Z	z
G3298u1	WIAF-15525	HT3504	1743 [		GGGCGGTGCT [G/A] CAGAACACGT	S		ᄓ	ᄓ
G3298u2	WIAF-13324	HT3504	2002		GTTCTTGCTG [A/G] AATGGTCCTT			×	凶
G3298u3	WIRE-13320	X82540	1044		AAGGCCAACA [C/T] AGCTGCAGGC			H	н
GSSMI.	WTAF-10255	X82540	1136	1136 INHBC, inhibin, beta C	CAGCAACATT [G/A] TCAAGACTGA			>	<u> </u>
G33113	WIAF-10256	X82540	1185	inhibin, beta C	GGGTGCAGTT [A/G] GTCTATGTGT	$\Box$	O E	* (	3 0
G33u4	WIAF-10259	X82540	892	a C	TTTTTGTGGA [C/T] TTCCGTGAGA	ν Ο		2	_إد
G3303u1	WIAF-13566	HT3523	981	POUGF1, POU domain, class 6, transcription factor 1	CAGGCCAGGA [G/A] ATCACTGAAA	လ ဂ	A	[22]	<u>FA</u>
G3304u1	WIAF-13932	HT3544	970	SP2, Sp2 transcription factor	TCAACAACCT [C/T] GTGAACGCCA	S	<u>H</u>	ㅁ	ы
G3304u2	WIAF-13935	HT3544	1891 SP2,	Sp2 transcription factor	AGAAGCACGT [T/G] TGCCACATCC	S	ט	<b>&gt;</b>	>
G3304u3	WIAF-13943	HT3544	920	SP2, Sp2 transcription factor	TGTGGTGAAG [T/C] TGACAGGTGG	S	U	리	ᆈ
2231111	WTAF-13839	HT3585	757	GATA3, GATA-binding protein 3	CCCACTCCCG [T/C] GGCAGCATGA	S	ני	ద	ద
6001100	WTAR-13840	HT3585	901	901 GATA3, GATA-binding protein 3	TCGGATGCAA [G/A] TCCAGGCCCA	S	A N	×	×
G3316u1	WIAF-13818	HT3607	282	zinc finger protein HKE-T1, Kruppel-like	AAAGAGTTTC[A/G]GTCAGAGTTC	Æ	Ö	Ø	_D
1,10	WT&R-14214	HT3613	1086	SMARCA3, SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 3	AAACTCTTAC [A/G] GCCAITGCAG	S	<u>v</u>	E-I	E+
				SMARCA3, SWI/SNF related, matrix associated, actin dependent requlator of chromatin, subfamily					
G3319u2	WIAF-14221	HT3613	1261	a, member 3	TAGATGTAGT [G/C] AACAACCCAG	Σ	ပ ပ	<b>X</b>	⊃ <b>'</b>
G3320u1	WIAF-13692	HT3622	624	BCL6, B-cell CLL/lymphoma 6 (zinc 624 finger protein 51)	ATTTGCGGGA [G/C] GGCAACATCA	Σ	ບ ຫ	E	

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G3320u2	WIAF-13717	HT3622	1062	BCL6, B-cell ( finger protein	51)	ACAGCCGGCC [G/A] ACTTTGGAGG	ω Ω	A	Д	д
				STAT2, signal activator of tr	transducer and canscription 2,					
G3321u1	WIAF-13761	HT3641	235	113kD		TCTTGGATCA [G/C] CTGAACTATG	<b>∑</b> .	ပ	2	티
				STAT2, signal activator of t	STAT2, signal transducer and activator of transcription 2,	Sabababacct [G/C] CATCAGAGCT	<u>U</u>		ບ	ω.
G3321u2	WIAF-13762	HT3641	4//	//4 LLSKU	factor znf6	CCACAATGGT [A/G] TCAGAGGAGG	S	Ö	>	>
G3328u1	WIAF-13543 WTAF-13544	HT3681	1389	1389 transcription	factor znf6	AGAGGATTTA [G/C] AGGAAGATGA	Æ	C)	田	0
			910	vod-X tody 2tc	x-how binding protein 1	ACCTGAGCCC [C/T] GAGGAGAAGG	Ω Ω	<del>[- </del>		വ
G3336u1	WIAF-13848	H15/32	26%			TACATTGGCC [A/C] CAAGACAAAG	MA	C	田	д
6334u1	WIAF-10000	1111220	2000		1	TCACAGCCCT [T/C] CGGCCAGGGT	M	נ	14	Ω
G334u2	WIAF-10009	H11220	1521		1	CCCAGATGAA [T/C] GGGAAACCCT	ST	ت ا	Z	Z
6334u3	WINE 10010	HT1220	2210		7	GGCTGGCCCA [A/G] TGAGAACCTG	M	O.	Z	S
6334 u4	WIRE-10019	HT1220	2979		1	GTGAGACCGA [T/C] TTCCGCCGAT				
2324116	WTAF-10033	HT1220	1136	1136 THBS1, throm	thrombospondin 1	TGTCACTGTC [A/G] GAACTCAGTT				
G334u7	WIAF-10034	HT1220	1859		thrombospondin 1	AGTGGAAATG [G/A] CATCCAGTGC	Σ	<b>₹</b>	U .	
G3343u1	WIAF-13545	HT3770	1104	ZNF76, zinc 1104 (expressed in	zinc finger protein 76 sed in testis)	GCAGTGCCCA [C/T] GGCGAGCTGG	ß	E-I	E.	田
2343112	MTA 7 2 7 2 7 6 1	HT3770	425	ZNF76, zinc finger p	zinc finger protein 76 ed in testis)	GAGCAGTATG [C/A] CAGCAAGGTT	Σ	C Y	A	<u> </u>
330				ZNF76,	zinc finger protein 76	, או איר איר איר איר איר איר איר איר איר איר	≥		<u>-</u> E-	<u>&gt;</u>
G3343u3	WIAF-13562	HT3770	143	(expressed in testis)	1 testis)	CACCAGGIGA [C/ 1] GGIACAGGIG				
G3343u4	WIAF-13563	HT3770	646	ZNF76, (express	zinc finger protein 76 ed in testis)	GAAGAGCCAC [G/T] TTCGTACCCA	Σ	ט	D I	<u>F</u>
23343115	WIAF-13564	HT3770	611	ZNF76, (express	zinc finger protein 76 ed in testis)	AGCTGTGGAA [A/G] GGCCTTTGCC				
G3344u1	WIAF-13664	HT3772	925	zinc finger	protein MAZ	AGCTGTCGCA [C/T] TCGGACGAGA	တ	บ	EH	H H
G3345u1	WIAF-13508	HT3823	315	TCF6L1, trans like 1 (mitoch 315 transcription	transcription factor 6- (mitochondrial iption factor 1-like)	TTCGATITTC [T/C] AAAGAACAAC	S	E	D D	ω ω

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2337311	WTAF-14203	HT4342	1253	MTF1, metal-regulatory	CTCAACAGAC [A/G] GCTTCCTTGA	8	ប	₽	
11106220	WTAF-14182	HT4483	680	ZNF133, zinc finger protein 133 (clone pHZ-13)	AGAGCCAGAG [C/T] TCTACCTCGA	υ Σ	E	그	Fa.
G3390112	WIAF-14184	HT4483	1026	ZNF133, zinc finger protein 133 (clone pHZ-13)	GCTCAGACAG [G/A] GAACCCTGAG	ڻ ع	A	_U	E
23390113	WTAF-14185	HT4483	1423	ZNF133, zinc finger protein 133 (clone pHZ-13)	AAAAGCCTTA [T/C] GTGTGCCGGG	co E-i	υ	N N	¥
71100000	WT&R-14197	HT4483	811	ZNF133, zinc finger protein 133 (clone pHZ-13)	CTGGGGATCC[A/G]GGCCCAGGGG	S	ტ	Д	Д
633390u#	WTAF-14198	HT4483	1420	ZNF133, zinc finger protein 133 (clone pHZ-13)	GGGAAAGCC [T/G] TATGTGTGCC	₽	U	Д	Д
9006885	WIAF-14199	HT4483	2143	ZNF133, zinc finger protein 133 (clone pHZ-13)	CAGCTCTAAT [C/T] ACACACAAGC	ນ ຮ	EH	н	н
G3391u1	WIAF-13631	HT4484	391	ZNF136, zinc finger protein 136 (clone pHZ-20)	AGCATTGTAT [A/G] TGGAGAAGTC	M	ტ	ы	υ
G3396111	WIAF-13978	HT4491	1283	ZNF135, zinc finger protein 135 (clone pHZ-17)	CACAGCTCCT [C/T] GCTCAGCCAG	υ Σ	<u>-</u>	ß	ы
G3396u2	WIAF-13979	HT4491	1296	ZNF135, zinc finger protein 135 (clone pHZ-17)	TCAGCCAGCA [C/T] GAAAGGACGC	S	E	д	ш
51130660	WTDF-13980	HT4491	1028	ZNF135, zinc finger protein 135 (clone pHZ-17)	AGTCACAGCT [C/T] GTCCCTCACC	υ Σ	_ ⊟	ß	ы
7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	MTATA 13981	HT4491	1057		GCGAATCCAC[A/G]CTGGGGAGAA	M A	υ	H	A
43396u5	WIAF-13982	HT4491	1152	ZNF135, zinc finger protein 135 (clone pHZ-17)	CAGGAGAGAA [A/G] CCCTATGAAT	S	U	×	×
91196282	WIAF-13983	HT4491	1243	ZNF135, zinc finger protein 135 (clone pHZ-17)	AAAGCCGTAT [G/C] GGTGCAATGA	ڻ ع	υ	ტ	D4
G3396u7	WIAF-13984	HT4491	1045	ZNF135, zinc finger protein 135 1045 (clone pHZ-17)	CACCAAACAT [C/T] AGCGAATCCA	z z	₽	α	*
G340u1	WIAF-10139	HT1386	459	CYP27A1, cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebrotendinous 459 xanthomatosis), polypeptide 1	cctatgggcc[G/A]ttcaccacgg	ν υ	<u> </u>	д	Д
G340u2	WIAF-10160	HT1386	801	CYP27A1, cytochrome P450, subfamily XXVIIA (steroid 27- hydroxylase, cerebrotendinous 801 xanthomatosis), polypeptide 1	TCCCCAAGTG [G/A] ACTCGCCCCG	z z		8	*

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2341111	WTAF-10121	HT1388	912 n	MUT, methylmalonyl Coenzyme A 912 mutase	GAGCTGGCCT [A/G] TACTTTAGCA	I A G	<u> </u>	Ü
G341u2	WIAF-10128	HT1388	2087	MUT, methylmalonyl Coenzyme A	TGCTGTGGGC [G/A] TAAGCACCCT M	٦ ٩	>	н
G3410u1	WIAF-13749	HT4550	1720 2	zinc finger homeodomain protein	TGAGTCCTCT [G/T] TTTCATCAGC	D E	>	Ĭτι
G3410u2	WIAF-13750	HT4550	2843	zinc finger homeodomain protein	AAACATCATT[T/C]GATTGAACAC	EH C	<u>п</u>	Ω
G3410u3	WIAF-13751	HT4550	2745	zinc finger homeodomain protein	AGATATTCCA [A/T] AAGAGTAGTT	A A	OX EH	н
G3410u4	WIAF-13775	HT4550	236	zinc finger homeodomain protein	AGAGAAGGGA [A/C] TGCTAAGAAC	Æ	υ V	H
G3410u5	WIAF-13776	HT4550	195	zinc finger homeodomain protein	TGCCAACAGA [C/T] CAGACAGTGT	S S	A H	
G3410u6	WIAF-13777	HT4550	909	zinc finger homeodomain protein	ADAACTITAG [1/C] TGCTCCCTGT	ω H	<u>ა</u>	တ
71011	WIAF-13793	HT4550	2073 zinc	finger homeodomain protein		A		Д (
G343u1	WIAF-10120	HT1552	561 HK1	, hexokinase 1		4 Z		۱ بح
G343u2	WIAF-10124	HT1552	159 HK1,	hexokinase 1	ACAAGTATCT [G/C] TATGCCATGC	Э	<u>ا د</u> ا د	4
G348u1	WIAF-10269	HT1906	2212	PECAM1, platelet/endothelial celladhesion molecule (CD31 antigen)	TGACGATGTC [A/G] GAAACCATGC	ه ه	<u>ი</u>	_O
G348u2	WIAF-10277	HT1906	1656	PECAM1, platelet/endothelial cell adhesion molecule (CD31 antigen)	GCCATTCCCA [C/T] GCCAAAATGT	υ w	H H	耳
2348113	WTAF-10283	HT1906	577	PECAM1, platelet/endothelial cell adhesion molecule (CD31 antigen)	AGAGTACCAG [C/G] TGTTGGTGGA	ე თ	۵ ک	<u>&gt;</u>
G3488 8885 8855 8855	WIAF-13119	9	٥.	PECAMI, platelet/endothelial celladhesion molecule (CD31 antigen)	ATTGTTCCC[C/G]	υ ~	U	
G351u1	WIAF-10123	HT1990	1047	1047 OSBP, oxysterol binding protein	TGCTGGCAGA [G/A] TCAGATGAAT	<u>ი</u>	A	田
G351u2	WIAF-10132	HT1990	1023	1023 OSBP, oxysterol binding protein	TGGCCAAGGC [C/A] AAAGCTGTGA			< ≥
G355u1	WIAF-10146	HT2143	1670	thrombospondin	AACTGCCTGA [G/A] TGTCTTAAAT	5 U	מ ע	ļ
G355u2	WIAF-10165	HT2143	1186	1186 THBS4, thrombospondin 4	TCGAAATGGA [G/C] CGTGCG11CC	٦.	7	1

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G355a3	WIAF-10510	HT2143	7967	1962 THBS4,	#			ď	⊧	-
G355a4	WIAF-13125	HT2143	1963	1963 THBS4,		CIGCCCACC [6/a] ICALIACAG		9 0	>  -	
G3552u1	WIAF-12701	HT28101	1006	1006 CLCN2,	chloride channel 2	AAGAGACTAT [T/C] ACAGCCCTCT		ا د	-1	-
G3552u2	WIAF-12731	HT28101	1823	1823 CLCN2,	chloride channel 2	CCGCCACCAG [C/T] AGTACCGGGT		H	ο	*
73552113	WTAF-12736	HT28101	2254	2254 CLCN2,	chloride channel 2	GGAGCGCAGA [G/C] TCGGCAGGCA	υ Σ	ပ	м	
G3565111	WTAF-12744	HT2896	334	334 calcyclin	-	GCCCTCAAGG [G/A] CTGAAAATAA	დ	∢	უ	Ω
G357u1	WIAF-10267	HT2244	4300 C4B,		plement component 4B	atgagtacga [t/c] gagcttccag	S	۷		
G357u2	WIAF-10280	HT2244	5095	C4B,	complement component 4B	TCATGGGTCT [G/A] GATGGGGCCA	S	4	ㅁ	귀
G357u3	WIAF-10295	HT2244	2996 C4B	C4B,	complement component 4B	CTCAGATCCA [T/C] TGGACACTTT	Ω [-ι	U		ᆈ
G359n1	WTAF-10026	HT2411	936	PLAT, tissue	plasminogen activator,	CGCAGGCTGA [A/G] GTGGGAGTAC	M	ŋ	E	Σ
1			777	PLAT,	plasminogen activator,	AGGCCTTGTC [T/C] CCTTTCTATT	S F	ັບ	<u> </u>	Ø
G359a2	WIAF-10520	HTZ411	744	743 CI,CN4	chloride channel 4	CITCTAACGA [G/A] ACCACITITG	S	A	M	ĮΧĮ
G3592U1	WIAF-12/37	HT4214	835	CLCN4,	chloride channel 4	GCTTACATTC [T/G] GAATTACTTA	E	Ü	ы	24
	WT&R-10053	НТ2479	857		cystathionine beta synthase, alt. transcript 1	TGGCTCACTA [C/T] GACACCACG	<u>ა</u>	F-		×
		0	000		cystathionine beta synthase, alt.	TCATCCCCAC [G/A] GTGCTGGACA	დ ტ	4	E	EH.
G361u2	WIAE-10056	0000E	COL	ADRB2,	ADRB2, adrenergic, beta-2-,	GGCACCAAT [G/A] GAAGCCATGC	_ უ	4	<u></u> ೮	ద
G362u1	W1AF-10058	H12038	677	ADRB2,	adrenergic, beta-2-,	בים חיבים חיבים ביים ביים ביים ביים ביים	ָ ני	_ A		-
G362u2	WIAF-10059	HT2638	429	receptor,	surface	TCATGGGCCT [6/A] GCAGTGGTGC	7		-	1
G362u3	WIAF-10060	HT2638	256	ADRB2, recept	ADRB2, adrenergic, beta-2-, 256 receptor, surface	CGTCACGCAG [G/C] AAAGGGACGA	Σ	ט	ш	a
6362114	WTAF-10093	HT2638	1230	ADRB2, a	adrenergic, beta-2-, or, surface	AGGCCTATGG [G/C] AATGGCTACT	S	D D	<u></u>	ŋ
G3620u1	WIAF-12808	HT97200	458	ACATN, ace 58 transporter	acetyl-Coenzyme A orter	CACTCTCTGG [A/G] TATGAAGAGC	Σ	<b>ʊ</b>	_	r.
G3627u1	WIAF-12820	HT97387	347	NAPG, factor	N-ethylmaleimide-sensitive attachment protein, gamma	GCAGAAACTA [C/T] CAGAGGCCGT	×	E D	<u>Р</u>	ß
G366u1	WIAF-10046	HT2764	987	987 BDKRB2,	, bradykinin receptor B2	GCCICCIICA [1/C] GGCCIACAGC	Σ	E E	Σ	H
G366a2	WIAF-10500	HT2764	820	820 BDKRB2	2, bradykinin receptor B2	AGATCCAGAC [G/A] GAGAGGAGGG	တ	G A	H	<u>-</u>

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G366a3	WIAF-105UL	H17710#	T D D	ACACA,					- 1	,
G367u1	WIAF-10156	HT27685	6965	carboxy	pha	ATCATCCATA [T/C] GACGCACC			* [	ء اد
G370u1	WIAF-10281	HT27888	3250	3250 LEPR,	leptin receptor	AAAATTCTCC [G/A] TTGAAGGATT		7	14 (	1, (
G370112	WIAF-10282	HT27888	3229	3229 LEPR,	leptin receptor	TCACCAAGTG [C/T] TTCTCTAGCA			ပ	راد
23.70113	WTAF-10284	HT27888	1005	1005 LEPR,	leptin receptor	CAATATCAAG [T/C] GAAATATTCA	E		>	4
C270114	WTAF-10285	HT27888	1894	1894 LEPR,	leptin redeptor	CAGAGAATAA [C/T] CTTCAATTCC	S C	EH	z	Z
63.70m	MTAE-10299	HT27888	1222	1222 LEPR.		TTCTGACAAG [T/C] GTTGGGTCTA	S)	O .	ß	S
G370116	WIAF-10300	HT27888	2161	2161 LEPR,	leptin receptor	CTATGAAAA [G/C] GAGAAAAATG	Σ	0	×	z
G371u1	WIAF-10107	HT27943	349	CRAI,	carnitine acetyltransferase	acetyltransferase TCATCTACTC[G/C]AGCCCAGGCG	თ	_ 0	_ ω	ω
G371a2	WIAF-12093	HT27943	287	CRAT,	carnitine acetyltransferase	GGAGAACTGG [C/T] TGTCTGAGTG	S	اط ن	ㅂ	ᆈ
				HADHA,	hydroxyacyl-Coenzyme A			_		
				dehydr	dehydrogenase/3-ketoacy1-Coenzyme					
				A thio	A thiolase/enoyl-Coenzyme A					
	1		7	hydratase (tr	hydratase (trifunctional procein),	TGGAGCTCCA [C/A] AGAAGGATGT	Σ	<u>ہ</u> ں	 	×
G372a1	WIAF-10506	UT2024	4435	4435 FASN	fatty acid synthase	CACCTCCCAC [G/A] TCCCGGAGGT	Σ	G A	V	н
63/4uT	MIAF - LOLOLO	00.000		MOKE		CTGGACAGGG [T/C] GACCCGAGAG	Σ	T C	>	Ø
G374u2	WIAF-10104	H128496	2880	Syyb FASIN,	מרדע	Caracarac (a) 41 Trancarac	Σ	A G	Н	Þ
G374u3	WIAF-10105	HT28496	5644	5644 FASIN,	acıu.	COPROCED FF/ CT BACKDACCER	1	E C	-	Н
G374u4	WIAF-10115	HT28496	6387	6387 FASN,	fatty acid synthase	1'GGCACACA1 [C/1] CIGGGCA1CC	1			$\top$
G374u5	WIAF-10119	HT28496	567	567 FASN,	fatty acid synthase	GGGGCATCAA [C/T] GTCCTGCTGA				
G374a6	WIAF-12094	HT28496	5520	520 FASN,	fatty acid synthase	ACATGGCCCA[A/G]GGGAAGCACA	တ	্চ ড	OX	≃ -
	2 c c c c c c c c c c c c c c c c c c c	900 CHn	929	PCCB, propi	propionyl Coenzyme A	GGACCCGGCT [T/C] CCGTCCGTGA	Σ	E E	ນ	ద
637/UL	WIAE-IOI44	HILOSO			T 47 T					
				PCCB,	propionyl Coenzyme A			<u></u>	, č	
G377u2	WIAF-10143	HT2996	1416	1416 carboxylase,	ylase, beta polypeptide	CACCTTTGTG [G/A] TGATACCAAC				
G380u1	WIAF-10122	HT3159	831	INSR,	insulin receptor	TCTACCTGGA [C/T] GGCAGGTGTG		T		
G380112	WTAF-10126	HT3159	1698	1698 INSR,	insulin receptor	GGCAGGATGC [A/G] TGTGGTTCCA				
G380u4	WIAF-11605	HT3159	2382	2382 INSR,	insulin receptor	GCGTGCCCAC [G/A] AGTCCGGAGG	ω	ر ن	T.	₽
				hdsohd	phospholipase C, beta 3, alt.		Σ		A R	0
G383n1	WIAF-10125	HT33546	363	3633 transcript 2	ript 2	AGCAGCGGGC [6/A] AGGC1CCCCC	-		Τ	T
		UT 2 2 8 3	ر 10	PRCP,	PRCP, prolylcarboxypeptidase	ATGACAGTGC [A/G] GGAAAGCAGC	S	Æ	ر لا	A .
G385U1	WIAT - IOL4I	n13303	2	- C						

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G385112	WIAF-10157	HT3383	1360	PRCP, (angio	PRCP, prolylcarboxypeptidase	ATCACAGACA[C/G]TCTGGTTGCA	υ	ڻ	Ð	ω
G387u1	WIAF-11729	HT3439	2697	SREBF2, 2697 binding	regulatory element ption factor 2	CACTCTCCAG [G/C] AGCTCCGTGC M	<b></b>	U	ĸ	Ø
	טרר דר מירוש סרר דר	н 13439	1901	SREBF2,	sterol regulatory element transcription factor 2	GCTGCTGCCG [C/G] CAACCTACAA	ບ	ū	4	rg.
G388u1	WIAF-110270	HT3440	245	245 SELPLG,	selectin P ligand	CTCCAGAAAT [G/A] CTGAGGAACA M	D _	Ø	Σ	н
G390u1	WIAF-10276	HT3568	2049	NOS3, (endot	NOS3, nitric oxide synthase 3 2049 (endothelial cell)	TTGCTCGTGC [C/G] GTGGACACAC	Ü	<u> </u>	<b>4</b>	
G391u1	WIAF-10013	HT3630	6205 VWF,	- 1	von Willebrand factor	AGGACCTGGA [G/C] GTGATTCTCC	υ U	υ	回	Д
G391u2	WIAF-10265	HT3630	4554	554 VWF,	von Willebrand factor	GCCCCTGAGA [A/G] CAAGGCCTTC M	A A	U	z	တ
G391u3	WIAF-10266	HT3630	7489	VWF,	von Willebrand factor	TGGCCTCAAC[C/T]GCCACCAATG	υ m	E	E	<u>-</u>
G391u4	WIAF-10272	HT3630	2470 VWF	VWF,	von Willebrand factor	ACTGTACCAT [G/A] AGTGGAGTCC	Ω.	Æ	Σ	н
G391u5	WIAF-10273	HT3630	2615 VWF	VWF,	von Willebrand factor	GCTCGAGTGT [A/G] CCAAAACGTG M	₹ _	ΰ	H	4
G391u6	WIAF-10274	HT3630	2635 VWF	VWF,	von Willebrand factor	GCCAGAACTA[T/C]GACCTGGAGT	EH S	U	<b>&gt;</b> -	Ħ
G391u7	WIAF-10275	HT3630	4045 VWF	VWF,	von Willebrand factor	TCTCGGAACC [G/A] CCGTTGCACG	o o	Æ		Ωı
G391u8	WIAF-10278	HT3630	4446 VWF	VWF,	von Willebrand factor	AACTTTGTCC [G/A] CTACGTCCAG	Σ Σ	Æ	r4	=
G391u9	WIAF-10279	HT3630	5152	VWF,	von Willebrand factor	GCCCTAATGC [C/T] AACGTGCAGG	ω U	E	A	Æ
G391u10	WIAF-10286	HT3630	3448	VWF,	von Willebrand factor	TTACCAGTGA [C/T] GTCTTCCAGG	S	E	Д	Д
G391u11	WIAF-10287	HT3630	4891	4891 VWF,	von Willebrand factor	ACATGGTGAC [C/T] GTGGAGTACC	S C	E4	E	E
G391u12	WIAF-10288	HT3630	4805	4805 VWF,	von Willebrand factor	CAGGAGCAAG [G/A] AGTTCATGGA	Σ U	4	[2]	ᅜ
G391u13	WIAF-10289	HT3630	4943	4943 VWF,	von Willebrand factor	CCTGCAGCGG [G/T] TGCGAGAGAT	∑ U	H	>	ᆈ
G391u14	WIAF-10290	HT3630	4915	4915 VWF,	von Willebrand factor	TCAGCGAGGC [A/C] CAGTCCAAAG	<b>α</b>	Ü	A	A

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G391a15	WIAF-10517	HT3630	6194 VWF,	ĺ	on Willek	von Willebrand factor	4	AAACAAGGAG [C/T] AGGACCTGGA	CTGGA	Z	Ü	E	0	*
G391a16	WIAF-13222	HT3630	6419 VWF		on Willek	von Willebrand factor	F	TCACCTTGGT [C/T] ACATCTTCAC	TTCAC	Σ	υ	E	н	<b>≯</b>
G3941u1	WIAF-14123	HT3464	1265	1265 mannosidase,		alpha, lysosomal	- O	CAGGTGTGCA [A/G] CCAGCTGGAG	TGGAG	Σ	Æ	<u>.</u> უ	Z	S
G3941u2	WIAF-14135	HT3464	965	965 mannosidase,		alpha, lysosomal	_ ₹	ACCAACCACA [C/T] TGTGATGACC	TGACC	Σ	บ		Н	н
G395u1	WIAF-10271	HT4158	1627	ECE1, 1627 enzyme	endotheli 1	endothelin converting	<u> </u>	TCACTGCCGA [T/C] CAGCTCAGGA	CAGGA	S	H	U	Д	Д
G395a2	WIAF-13110	HT4158	1493	ECE1, 1493 enzyme	endothelin 1	n converting	3	CATCTACAAC [A/T] TGATAGGATA	GGATA	Σ	Æ	E+	Σ	ы
G3959u1	WIAF-13634	HT4490	250	ADTB1, 250 prime)	adaptin,	beta 1 (beta	F-	TGAAGAAGCT [G/A] GTATACCTCT	CCTCT	w	U	Æ	1	F.J
G3959u2	WIAF-13640	HT4490	2029	ADTB1, 2029 prime)	adaptin,	beta 1 (beta	H	TTCTTGGCGG [T/C] GGCCTTGACA	TGACA	S	E	υ	ט	U
G3959u3	WIAF-13641	HT4490	2395	ADTB1, 2395 prime)	adaptin,	beta 1 (beta	4	AGGTCCACGC [G/A] CCACTCAGCC	CAGCC	ഗ	ප	Æ	Æ	Æ
G3967u1	WIAF-13997	HT2958	918	ACTC, 918 muscle	actin, al	alpha, cardiac		GAGGCACCAC [1/C] ATGTACCCTG	CCCTG	S	H	υ		E
G3968u1	WIAF-14159	HT1986	1747	1747 ACTN3,	actinin,	alpha 3		CGAGGCTGAC [C/T] GAGAGCGAGG		z	U	E	ద	*
G3968u2	WIAF-14164	HT1986	1900	1900 ACTN3,	actinin,	alpha 3	G	GGTGCCCAGC [C/T] GTGACCAGAC		Σ	U	E H	PH.	U
G3968u3	WIAF-14165	HT1986	2184	2184 ACTN3,	actinin,	alpha 3	A	ACACCGTCTA [C/T] AGCATGGAGC	GGAGC	S	C	T	Ϋ́	Y
G3968u4	WIAF-14167	HT1986	2557	2557 ACTN3,	actinin,	alpha 3	Ŋ	GATCTTGGCA [G/A] GAGACAAGAA		Σ	G	A (	G	R
G3968n5	WIAF-14175	HT1986	1212	1212 ACTN3,	actinin,	alpha 3	D	GGCTGCTCTC [G/A] GAGATCCGGC	၁၅၅၁၁	S	9	Ą	ß	S
G3979u1	WIAF-13884	HT0623	176	776 GPC1,	glypican	1	T	TGCTGCTGCC [T/G] GATGACTACC	CTACC	လ	E	ro O	д	Д
G3979u2	WIAF-13885	HT0623	680	680 GPC1,	glypican	1	I	TGTACTACCG [C/T] GGTGCCAACC	CAACC	വ	ນ	E	24	ద
G3979u3	WIAF-13886	HT0623	1361	1361 GPC1,	glypican	1	A	AGCTGGTCTC [T/C] GAAGCCAAGG	CAAGG	ω	E	ບ	Ω.	တ
G3979u4	WIAF-13887	HT0623	1163	1163 GPC1,	glypican 1	1	A	AGAGTGTCAT [C/T] GGCAGCGTGC	CGTGC	S	ນ	ы	I	н
G3979u5	WIAF-13888	HT0623	1670	1670 GPC1,	glypican	1	4	ACGCCAGTGA [C/T] GACGGCAGCG	CAGCG	S	บ	ī	Q	Д
G3979n6	WIAF-13905	HT0623	1069	1069 GPC1,	glypican	1	J	CTTGCCAACC [A/T] GGCCGACCTG		Σ	A	E	α	П
G3979u7	WIAF-13906	HT0623	1514	1514 GPC1,	glypican	1	T	TCATGGGTGA [C/T] GGCCTGGCCA	GGCCA	ß	ນ		D	D
G3979u8	WIAF-13907	HT0623	1720	1720 GPC1,	glypican 1	F	9	GACCTCTGCG [G/C] CCGGAAGGTC		Σ	U	ں ت	ט	Æ
G3979u9	WIAF-13908	HT0623	1676	1676 GPC1,	glypican	1	ъ	GTGACGACGG [C/T] AGCGGCTCGG	CICGG	S	U	E	ט	U
G3979u10	WIAF-13909	HT0623	1719	1719 GPC1,	glypican	1	T	TGACCTCTGC [G/A] GCCGGAAGGT		Σ	ט	A	ט	ς Ω
G399u1	WIAF-10102	HT48511	450	450 AQP3,	aquaporin	1 3	T	TCTGGCACTT [T/C] GCCGACAACC	CAACC	Ω	E	ט	[24	ſ±ι
G399u2	WIAF-10111	HT48511	192	192 AQP3,	aquaporin 3	1 3	D	GCTCCGTGGC [C/T] CAGGTTGTGC	TGTGC	ω	ບ	 H	Æ	Æ
G399u3	WIAF-10112	HT48511	165	165 AQP3,	aquaporin	1 3	U	CCCTCATCCT [C/G] GTGATGTTTG	GITIG	ω	U	ט	ı	ıп
G3997u1	WIAF-13649	HT27682	473	MFAP2, 473 protein		microfibrillar-associated 2		TGTGTGCCCA [C/T] GAGGAGCTCC	GCTCC	Ω	υ	٤	H	щ
G3997u2	WIAF-13650	HT27682	377	MFAP2, 377 protein		microfibrillar-associated 2		CCATACACAG [G/T] CCTTGCAAAC		Σ	G	T.	PK.	S
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G3997u3	WIAF-13876	HT27682	453	MFAP2, microfibrillar-associated 453 protein 2	GGAGATCTGT [G/T] TTCGTACAGT	ט	- A	Eta
G4022u1	WIAF-14020	HT2426	240 9	TGM1, transglutaminase 1 (K polypeptide epidermal type I, protein-glutamine-gamma- glutamyltransferase)	TGGCTGCTGT [T/C] CATGCCGAAA	E	ა ე	<u>Δ</u> ,
G4022u2	WIAF-14021	HT2426	371	TGM1, transglutaminase 1 (K polypeptide epidermal type I, protein-glutamine-gamma- glutamyltransferase)	CCCGGGGCAG[C/T]GCTGTCAATG	<u>ი</u>	E E	တ
G4 0 2 2 u 3	WIAF-14022	HT2426	506	TGM1, transglutaminase 1 (K polypeptide epidermal type I, protein-glutamine-gamma-glutamyltransferase)	ACGAGCTGAT [A/G] GTGCGCCGCG	M A	р Н	Σ
G4022u4	WIAF-14031	HT2426	2491	TGM1, transglutaminase 1 (K polypeptide epidermal type I, protein-glutamine-gamma-glutamyltransferase)	GCTGGAGGTG [A/T] CAGTCACTTA	Æ E	T D	> >
G4038u1	WIAF-13998	HT4211	411	<pre>LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))</pre>	GGTGGCAGTC[C/A]CAGAATGATG	ე ა	δ 2	ω
G4038u2	WIAF-13999	HT4211	258	<pre>LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))</pre>	CTTCATCTAC[C/T]TGTGGACTGA	ე დ	E E	E-1
G4038113	WIAF-14002	HT4211	1830	<pre>LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))</pre>	GAGGCTACTG [C/T] AATCGCTACC	<u>ი</u>	ن ا	ט
G4038u4	WIAF-14003	HT4211	2668	<pre>LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))</pre>	GACCAGGCAG [A/T] TGATTAGGGC	æ	E E	디
G4038u5	WIAF-14018	HT4211	248	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))	TITCTCCGAG [C/T] TTCATCTACC	υ Σ	H H	A \
G4038u6	WIAF-14019	HT4211	887	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))	CACGGCCATG [C/T] TGATCGCTGC	υ Σ	E	A V
G4038u7	WIAF-14023	HT4211	1266	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))	AGTGTGATCC [G/A] GATGGGGCAG	හ ව	A	
G4038u8	WIAF-14025	HT4211	1693	<pre>LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))</pre>	CTATGGAGAC[G/A]TGGCCACAGG	<u>ی</u>	Æ	×

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G4038u9	WIAF-14026	HT4211	1553	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))	GGCTGTGAAC [C/T] GTGTGCCTGC	Σ	E U	д	ᅱ	
G4038u10	WIAF-14029	HT4211	3562	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD))	CCTGACAGGA [C/T] TGGAGAAGCG	S	H ن		니	
	, tre Fra	1 C 1 C 1 T L	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	LAMB3, laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (195kD),	TGCTGCGCTC [A/G] GCGGACCTGA	ß	<u></u>	<u> </u>	თ	
G4035ULL	WTAF-13571	HT0652	1266	adducin, beta subunit	TGGAGCAGGA [G/T] AAGCACCGGC		E E	N	Д	
G4050u1	WIAF-14106	HT1466	1366	1	CGTTTGGCAG [G/A] GCAGCCAGGC	M	G A	ש	ഗ	
G4050u2	WIAF-14107	HT1466	1468	1468 villin	GGTCCCAATG [G/A] GCAAGGAGCC	Σ	<u>ه</u>			Т
G4050u3	WIAF-14108	HT1466	1932	1932 villin	CCACAGAGAT [C/T] CCTGACTTCA				1	Т
G4050u4	WIAF-14110	HT1466	2438	2438 villin	TTTGGGATGA [C/T] TCCAGCTGCC					Т
G4057u1	WIAF-13648	HT33633	371	371 CNN3, calponin 3, acidic	TTCAGGCTTA [T/C] GGTATGAAGC					Т
G4066ul	WIAF-13676	HT4301	654	654 troponin T, beta, skeletal	AGATTGACAA [G/A] TTCGAGTTTG					Т
G4066u2	WIAF-13677	HT4301	774	774 troponin T, beta, skeletal	GCAAAGTCGG [C/T] GGGCGCTGGA			_		Т
G4066u3	WIAF-13708	HT4301	625	625 troponin T, beta, skeletal	GGAGCTCTGG [G/C] AGACCCTGCA	Σ	ט	C)	의	Т
G4080111	WIAF-14142	HT1396	13130	HSPG2, heparan sulfate 13130 proteoglycan 2 (perlecan)	GATTCTCCTC [G/A] GGCATCACAG	လ	U	<b>₹</b>	ß	
G4080u2	WIAF-14150	HT1396	10340	10340 proteoglycan 2 (perlecan)	TTGAGTTCCA [C/T] TGTGCTGTGC	ω.	ار	ਸ਼  ∷-	디	T
G4080u3	WIAF-14151	HT1396	12392	HSPG2, heparan sulfate proteoglycan 2 (perlecan)	AATGCTATGA [T/C] AGCTCCCCAT	တ	H	A U	<u> </u>	
200	74 TAT 20 C Z 1 A 1 C 2	HT1 2 9 K	3416	HSPG2, heparan sulfate	TGGCTGTGCC [C/T] GAGGAAACCG	ß	 ບ	<u></u>	<u> </u>	
£2000£5	10111 TETH			1 -					_	
G4080u5	WIAF-14154	HT1396	4588	glycan 2 (	GTGCCGCTGG [T/C] GGCCAGCATC	×	E-1	D C	A.	
	L T	000	0 10	HSPG2, heparan sulfate	GGACACCAC [G/A] CGGIGCIGCA	Σ	ט	A A	[4	
G408046	WIAF-14130	HT4237	394	1	CAAAGAAATC [G/A] ATTCAGTCGG	ß	ט	A S	S	
G4096u2	WIAF-13910	HT4237	455	455 motor protein	ATCTAAACAG [C/T] CTGCCTCACA	Σ	Ü	ഥ		
G4096u3	WIAF-13911	HT4237	1150	1150 motor protein	CTAAGGTTGT [A/G] TCTCAGTATC	လ	Æ	D C	Δ .	T
G4109u1	WIAF-14034	HT28223	1238	phosphoglucomutase-related protein TACAGCGTGG[C/T]GAAGACGGAT	TACAGCGTGG [C/T] GAAGACGGAT	Σ	υ	- L	A A	Ţ
G4109u2	WIAF-14035	HT28223	1043	1043 phosphoglucomutase-related protein ATTATTGCTG[C/A]CCGGAAGCAG	ATTATTGCTG[C/A]CCGGAAGCAG	Σ	Ü	A	A.	
G4112u1	WIAF-13615	HT4401	374	KIF5A, kinesin family member 5A	AGATGTCCTT [G/A] CTGGCTACAA	Σ	b	A	A A	E
G4112u2	WIAF-13623	HT4401	2767	2767 KIF5A, kinesin family member 5A	AGAGAGTTAA [G/T] GCCCTGGAGG	Σ	ט	- H	M	z

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G4114u1	WIAF-14113	HT4160	830 1	, light polypeptide			, F		k
G4118u1	WIAF-14010	HT0841	564 5	mlatory	TCGATGTGGC [6/A] GGCAACCIGG		¢	¢	¢
G4118112	WIAF-14011	HT0841	368	MYL5, myosin, light polypeptide 368 5, regulatory	TTCACCATGT [T/C] TCTGAACCTG M	H	υ	[It ₄	ß
	CLOAL TATE	HT0841	7 2 2 2	, light polypeptide	GAGGIGGACC [A/G] GATGITCCAG	Æ	უ	Ø	R
G4112n1	WIAF-13955	HT97538	161 11	sin-I		SA	ນ	ı	ᆈ
G4124u1	WIAF-13895	HT0925	1517 9	TGM3, transglutaminase 3 (E polypeptide, protein-glutamine- gamma-glutamyltransferase)	TCGCTGGCAT [G/A] CTGGCAGTAG	Σ Ω	A	Σ	н
		L	0000	TGM3, transglutaminase 3 (E polypeptide, protein-glutamine-	AACCCAACAC [G/A] CCATTTGCCG	ა დ	Æ	단	E
G4124u2	WIAF-13896	10001H	10201	1039 myosin binding profein H		ນ	ט	S	S
G4126U1	WLAF - 13050	1112405	2007	369 myosin binding protein H		_ເ	ນ	Ö	Æ
G4.126.02	WIAF-13033	HT1657	198			⊡	υ	Д	Д
100000000000000000000000000000000000000	WTAR-13598	HT33664	601	MAGP2: Microfibril-associated glycoprotein-2	GAAAGATGAG [C/T] TTTGCCGTCA	υ Σ	E	ᆸ	[tı
011000	WIAH-13599	HT33664	405	MAGP2: Microfibril-associated	ATGACTTGGC [C/T] TCCCTCAGTG	ر د	E	Ą	Æ
G4138u3	WIAF-13600	HT33664	327	MAGP2: Microfibril-associated 327 glycoprotein-2	AAGATCCTAA [T/C] CTGGTGAATG	E S	U .	z	z
1,027	MTAR-14048	HT3443	1119	SNL, singed (Drosophila)-like (sea urchin fascin homolog like)	GCTGCTACTT[T/C]GACATCGAGT	ω -	U	ᄕᅭ	Ĺτι
Z4170m1	WTAF-13580	HT5069	1131	Golgi protein, peripheral, 1131 brefeldin A-sensitive	GAAATATACC [A/G] TAAGTATGGA	M A	Ü	н	>
2417002	WTAF-13581	HT5069	930	Golgi protein, peripheral, 930 brefeldin A-sensitive	GTATAATAAA [C/T] TCCTGGAGTT	υ Σ	E	ᆸ	Fz.
G4170u3	WIAF-13582	HT5069	2312	Golgi protein, peripheral, 2312 brefeldin A-sensitive	AGCAGCCTTA [A/G] GCATCTTGGA	N A	២	*	*
G41 70"4	WTAF-13596	HT5069	359	Golgi protein, peripheral, 359 brefeldin A-sensitive	TCAACCAGCT [T/G] TCTGTGCCTT	Ω ⊟	Ö	卢	니
G4170u5	WIAF-13597	HT5069	1007	Golgi protein, peripheral, 1007 brefeldin A-sensitive	AAAAAGGCAA [T/A] ACTGTTCCTG	E Z	4	z	
G4171u1	WIAF-13688	HT1587	667	667 KIFSB, kinesin family member 5B	TTTTAATTA[T/C]ATTTACTCCA	S F-I	Ü	<u>&gt;</u>	_×

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64171112	WIAF-13689	HT1587	1036	1036 KIF5B, kinesin family member 5B	TTAGTAAAAC [T/C] GGAGCTGAAG	S H	Ü	E E
G4176111	WIRF-14204	HT33754	130	TNR, tenascin R (restrictin, janusin)	GCTCATTGGC [G/A] TCAACCTGAT	υ Σ	Æ	H >
G4176u2	WIAF-14205	HT33754	463	TWR, tenascin R (restrictin, janusin)	CTGTCCATGT [G/T] CCAGTTCAGC	<u>ත</u>	E	.α .α
64176113	WTAF-14206	HT33754	249	TNR, tenascin R (restrictin, 249 janusin)	ACTACAACAC [G/A] TCCAGCAAAG	ა ტ	Æ	E E
G4176u4	WIAF-14208	HT33754	2009	TMR, tenascin R (restrictin, 2009 janusin)	CTGGTCCCCA [G/A] GGGCATTGGT	υ Σ	Æ	ద
G4176u5	WIAF-14209	HT33754	2175	TNR, tenascin R (restrictin, janusin)	CAGCCTCCTC [G/A] GAGACCTCCA	დ დ	Æ	S)
241 76116	WIAF-14210	HT33754	3318	TNR, tenascin R (restrictin, janusin)	AATCCACCGA [C/T] GGAAGCCGCA	လ လ	H	<u> </u>
24176117	WT AF-14211	HT33754	3221	TNR, tenascin R (restrictin, 3221 janusin)	CCGGCAAACC [T/C] GACAGCCAGT	E E	υ	ഥ
0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	WTAR-14217	HT33754	1635	TNR, tenascin R (restrictin,	TCTCGGACAC [C/T] GTGGCTTTTG	ນ	E	E E
G4178u1	WIAF-14138	HT0224	2827	2827 ACTN2, actinin, alpha 2	GCTGCGTTCT [C/T] TTCCGCACTC		E	
G4178u2	WIAF-14139	HT0224	2818	2818 ACTN2, actinin, alpha 2	CTGGATTACG [C/T] TGCGTTCTCT	υ Σ	ы	A
G418u1	WIAF-11750	L07594	2370	TGFBR3, transforming growth factor, beta receptor III (betaglycan, 300kD)	GAGTGCACTT [C/T] CCTATCCCGC	<u>ပ</u>	E	E4
641 8112	WIAF-11751	107594	25.86	TGFBR3, transforming growth factor, beta receptor III (betaglycan, 300kD)	AGAAGACGTT[C/T]ACCAAGCCCC	<u>ن</u> 2	E	[74 [74
G418u3	WIAF-11752	107594	2671		AATTTCTCCA [C/T] CAATTTTCCA	υ Σ	E	ъ О
G418u4	WIAF-11771	L07594	4.38	TGFBR3, transforming growth factor, beta receptor III (betaglycan, 300kD)	TGTGTGAACT [G/T] TCACCTGTCA	ν υ	E	<u>н</u>
G418u5	WIAF-11744	1.07594	392	TGFBR3, transforming growth   factor, beta receptor III   1922 (betaglycan, 300kD)	CTGATGAGCT [T/C] CTGTTTAGCC	Ε Σ	υ	ĺ¥ι

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G418u7 WIAF-11773 G418u8 WIAF-11745 G418u1 WIAF-11746 G4181u1 WIAF-14207 G4181u3 WIAF-14218 G4181u4 WIAF-14219 G4181u5 WIAF-14219 G4181u5 WIAF-14207 G4181u5 WIAF-14207	L07594 L07594 L07594 HT2008	1170 ( 11463 ( 12211 ( 12511 ( 12511 (	TGFBR3, transforming growth factor, beta receptor III  1170 (betaglycan, 300kD)  TGFBR3, transforming growth factor, beta receptor III  1463 (betaglycan, 300kD)	TCTTGAAGTG [C/A] AAAAAGTCTG				п
11 22 23 11		1463 ( 12211 ( 2211 ( 225 (	transforming growth beta receptor III ycan, 300kD)		υ z	Æ	υ U	*
11 12 23 11		2211 ( S		CCTCCTGAGC [T/C] ACGGATCCTG	Ε	υ	П	ρı
		425 6	TGFBR3, transforming growth factor, beta receptor III (betaglycan, 300kD)	ATGTTGAGGT [A/G] TCTGTTACTA	<u>م</u>	ט	>	>
			SPTBN1, spectrin, beta, non-	CTCTGCGCGG [C/T] TTTTTGAGCG	υ Σ	E	П	[Iza
	HT2008	3565	SPTBN1, spectrin, beta, non-	AGACAGCGAT [C/T]GCCTCGGAGG	ت ھ	EH	н	н
		1258	SPTBN1, spectrin, beta, non- erythrocytic 1	ACCTTCTGGA [A/G] TGGATTGAAC	ري دي	_O	M	凶
		1780	SPTBN1, spectrin, beta, non-	AGCTCGAGGC[C/T]GAGAATTACC	ى ت	₽	Æ	Æ
		3637	trin, beta, non- 1	acatcaagaa [t/c]gagatcgaca		U		z
		404	nyosin 4	CCAAGCACAT [T/C] GCGGAAGAGG	S	υ	н	ы
G418511   WIAF-13554		257	MFAP1, microfibrillar-associated protein 1	AAGGCCAGAC [T/G] ATGCCCCTAT	E E	G	¥	Д
		1108	MFAP1, microfibrillar-associated	CCAACAAAGC [T/G] GTTAAGGGCA	Ω [∺	Ŋ	Æ	A
		274	microfibrillar-associated	CTATGGAGTC [C/T] TCAGATGAGG		E-I	ß	S
		941	nucleoporin 88kD	GGGTCCATTG [C/A] CCATGCATCT		A	<b>⊲</b> 1	
		1092	, nucleoporin 88kD	ATGACCACAC [G/A] TCAGAAAGT		<b>₫</b>	-	٦ ,
	7 HT97558	1551	nucleoporin	TCCATCCAGC [G/A] TCTCCTCCCC		4 7	4 D	4 Þ
G4196u4 WIAF-13668		2220	nucleoporin	AGGGTGAACA [T/C] ATAAGGGAAA	ט מ	ט פ	<b>d</b> ≥	<b>□</b> ×
G4196u5 WIAF-13669		2205	ω	CCAICCIGAA [A/ G] GAGGAGGGIG		, 0	( E	: 0
		1329 VCL,	VCL, vinculin	CCATCTCCCC [A/G] ATGGTGATGG		ט	Д	, д
G4208u2 WIAF-13922	4 HT1122	818		GGGATGAAGA[T/C]GCCTGGGCCA	S	υ	Ω	Д
		1556 VCL,		AAGCACAGCG [G/A] TGGATTGATA	S D	K	ద	ద

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G4213u1	WIAF-13605	HT2813	163	163 NUP153,	nucleoporin 153kD	GCCAGGGTGG [T/C] TACAAAGATA	S	נ	ı	Г
G4213u2	WIAF-13606	HT2813	742	742 NUP153,	nucleoporin 153kD	GAATTCTTCA [A/G] TCCTTAAAAC	MA	r.	н	Ν
G4213u3	WIAF-13609	HT2813	1800	1800 NUP153,	nucleoporin 153kD	TTAGACCTGC [A/C] GAAATCCTGA	SA	U 1	ď	Æ
G4213u4	WIAF-13627	HT2813	1829	1829 NUP153,	nucleoporin 153kD	AGTGTTCTAG [A/C] TATTCTGAAA	Æ	ט	<u> </u>	Ø
G4213u5	WIAF-13632	HT2813	3258	3258 NUP153,	nucleoporin 153kD	CTTTTGGCAA [C/T] GTGGAGCCTG	S C	H	z	z
G4213u6	WIAF-13635	HT2813	4162	4162 NUP153,	nucleoporin 153kD	CTCTGGAACA [A/G] CTCCTAATTC	Σ	ט	H	Æ
G4218u1	WIAF-13854	HT1681	1122	phosphat 1122 class A	phosphatidyl-inositol glycan, class A	AACCTTATTA [T/C] TTTATGTGAG	Σ	T C	H	⊢
				CD36L2,	CD36L2, CD36 antigen (collagen					
G4223111	WIAF-14160	HT1684	1434	type i rereceptor)	type i receptor, intomosponari receptor)-like 2 (lysosomal integral membrane protein II)	ATTAGATGAC [T/C] TTGTTGAAAC	×	U E	[Yu	ᄓ
				CD36L2, CD36 a type I receptor receptor)-like	CD36 antigen (collagen receptor, thrombospondin or)-like 2 (lysosomal					
G4223u2	WIAF-14173	HT1684	969	integral	integral membrane protein II)	GTGGTCCCAG [G/A] TGCACTTCCT	Σ	<b>ধ</b> ড	>	Ξ.
G4223u3	WIAF-14174	HT1684	986		CD36L2, CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 2 (lysosomal integral membrane protein II)	CAGACAAGIG[C/I]AAIATGAITA	Ø	H ت	υ	ŭ
				CD36T.2	ch36 antigen (collagen					
	_			type I r	receptor, thrombospondin					
				receptor) -like	)-like 2 (lysosomal	ביבים אי איבוח ביי, בי חחות אבח אב יי			- 1	F
G4223u4	WIAF-14176	HT1684	1437	integral		AGATGACTTT [G/A] ITGAAACGGG		בן א ט כ		1 2
G4227u1	WIAF-14056	HT1929	912	proteoglycan	ycan 2	AIGCCICCAA [6/A] AAAGAIGGGG	o co			4 4
G4227u3	WIAF-14058	HT1929	1321	proteoglycan		CCGAGGAGC [T/C] ACTGGCGTCG		"		H
		C C	ī	SDC4, Sy	syndecan 4 (amphiglycan,	2504450440417/4140405450405	Σ		[z.	
G4229u1	WIAF-13901	HT4995	602	602 TRAM protein	tein	CCATAACCTG [A/C] TGACATTTCA		$\top$	Γ	П
G4243u1	WIAF-14169	HT2901	406	406 KRT17,	keratin 17	AGCTGGAGGT [G/A] AAGATCCGTG	ß	G B		>
G4243u2	WIAF-14170	HT2901	478	478 KRT17,	keratin 17	ACAGGACAAT [T/C] GAGGAGCTGC	ω	T C	H	н
G4243u3	WIAF-14171	HT2901	389	389 KRT17,	keratin 17	GGAGGAGGCC [A/G] ACACTGAGCT	Σ	A G	Z	
G4243u4	WIAF-14178	HT2901	564	564 KRT17,	keratin 17	CIGGCIGCIG [A/C] IGACITICGGC	Σ	A	C C	Æ
G4244u1	WIAF-14086	HT1056	386	clathrin	386 clathrin, light polypeptide a	ATCGATTGCA [G/C] TCAGAGCCTG	М	ro l	ر د	H
G4246u1	WIAF-14044	HT97492	259	259 SLN, sa	sarcolipin	GTCCTATCAG [T/C] ACTGAGAGGC			7	
G4246u2	WIAF-14045	HT97492	189	189 SIN, sa	sarcolipin	ACACCCGGGA [G/A] CTGTTTCTCA	S	G G	<u>H</u>	四

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G4254u1	WIAF-13546	HT3393	86 TNNIZ,	I2, troponin I,	skeletal, fast	ACCTGAAGAG [C/T] GTGATGCTGC	ຮ	E→	Ω	S
G4254u2	WIAF-13553	HT3393	530 TNNI2	I2, troponin I,	skeletal, fast	TCGAGGAGAA [G/C] TCTGGCATGG	<b>∑</b>	υ	×	z
G4255u1	WIAF-13644	HT2907	562 CRYAB	AB, crystallin,	alpha B	AGTTCCACAG [G/A] AAATACCGGA	ω U	A.	PK PK	p4
G4255u2	WIAF-13645	HT2907	367 CRYAB,	AB, crystallin,	alpha B	CCTCCTTCCT[G/A] CGGGCACCCA	ω υ	4	н	H
G4255u3	WIAF-13872	HT2907	271 CRYAB,	AB, crystallin,	alpha B	CCAGCCGCCT [C/T] TTTGACCAGT	ຽ	H	니	ᆈ
G4255u4	WIAF-13873	HT2907	580 CRYAB,	AB, crystallin,	alpha B	GGATCCCAGC [T/C]GATGTAGACC	ω	Ü	A	Æ
G4257u1	WIAF-14052	HT1694	PIGF, 394 glyca	PIGF, phosphatidylinositol glycan, class F	inositol	TAGAGTTGGC [A/G] TTGGAAACAT	S	<u></u> 5	A.	A
G4257u2	WIAF-14053	HT1694	PIGF, 252 glyca	PIGF, phosphatidylinositol glycan, class F		TATTTAGTAG [T/C] GAAACCAAAT	E	υ -	>	Æ
G4257u3	WIAF-14069	HT1694	PIGF, 291 glyca	PIGF, phosphatidylinositol glycan, class F		TCATTAICAC [A/G] CAAGGIAACT	Æ E	U U	田	<b>K</b>
G4264u1	WIAF-13519	T0968	TJP1,	<ol> <li>tight junction protein na occludens 1)</li> </ol>	on protein 1	CGGTCAGTGG [C/T] TTCCAGCCAG	Σ	H ن	Æ	>
G4264u2	WIAF-13520	HT0968 2:	TJP1, 2272 (zona	°	tight junction protein 1	CATGCTGATG [A/G] TCACACACCT	Σ	A G	D	ט
G4264u3	WIAE-13529	HT0968 5-	TJP1, 5408 (zona	1, tight junction protein ma occludens 1)	on protein 1	AGCCTCCTGA [A/T] GCTGATGGTG	Σ	A.	田	<u> </u>
G434u1	WIAF-11748	M21121	SCY 286 A5	SCYA5, small inducible 286 A5 (RANTES)	ible cytokine	TACATCAACT [C/T] TTTGGAGATG	Σ	C	Ω	[Eq.
G434u2	WIAF-11749	M21121	SCY 137 A5	SCYA5, small induc A5 (RANTES)	inducible cytokine	GCTTTGCCTA [C/T] ATTGCCCGCC	ω	H ن	>1	
G435u1	WIAF-11741	M31933	FCG 754 aff	FCGR2B, Fc fragmen affinity IIb, recep	Fc fragment of IgG, low IIb, receptor for (CD32)	GTCACTGGGA [T/C] TGCTGTAGCG	Σ	E-	H .	F
G435u2	WIAF-11743	M31933	FCGR2B, 395 affinity	FCGR2B, Fc fragment of affinity IIb, receptor	nt of IgG, low	GGGAGTACAC [G/A] TGCCAGACTG	ω	۵ 4	<u> </u>	H

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G435u3	WIAF-11742	M31933	673 年	FCGR2B, Fc fragment of 1gG, low 673 affinity IIb, receptor for (CD32)	TACACGCTGT [T/A] CTCATCCAAG	E E	4	[Et ₁	×
G4369u1	WIAF-13728	HT0900	G B B B D D D	GBE1, glucan (1,4-alpha-), branching enzyme 1 (glycogen branching enzyme, Andersen disease, glycogen storage disease	TTACGTCCAT [G/A] CTTTATCATC	<u>ა</u>	Æ	Σ	н
G4369u2	WIAF-13729	HT0900	9 b b	GBE1, glucan (1,4-alpha-), branching enzyme 1 (glycogen branching enzyme, Andersen disease, glycogen storage disease 1609 type IV)	GAGTCCTG [A/G] CTCCTTTAC	<b>ب</b> ح	_©	E	A
G4373u1	WIAF-13559	HT0940	1117 d	HSD17B2, hydroxysteroid (17-beta)	GCCAGCAAGG[A/T]CTTCTCTCCG	Æ	H	Ω	Λ
G4373u2	WIAF-13560	HT0940	1195 d	HSD17B2, hydroxysteroid (17-beta)	CCAGGGAAAG [G/A] CGCTTACTTG	D M	A	_O	Д
G438u1	WIAF-11830	M63121	583 T	TNFRSF1A, tumor necrosis factor 583 receptor superfamily, member 1A	ACCGTGTGTG [G/A] CTGCAGGAAG		Æ	ט	Д
G438u2	WIAF-11790	M63121	618 1	TNFRSF1A, tumor necrosis factor 618 receptor superfamily, member 1A	TTATTGGAGT [G/A] AAAACCTTTT	ტ <b>Σ</b>	Æ	Ħ	M
G440u1	WIAF-11806	M74447	Z61 b	TAP2, transporter 2, ABC (ATP 261 binding cassette)	TGCTAAAGCT [A/G]AGAGGGCTGC	ଓ ୟ	ტ	ы	ᄓ
G440u2	WIAF-11807	M74447	Z 2089 b	TAP2, transporter 2, ABC (ATP 2089 binding cassette)	CAGGCTGCAG [G/A] CAGTTCAGCG	<u>ت</u>		Æ	E⊣
G440u3	WIAF-11808	M74447	2155 k	TAP2, transporter 2, ABC (ATP 2155 binding cassette)	TGCCCAGCTC [C/T] AGGAGGGACA	ν υ	E	α	*
G440u4	WIAF-11818	M74447	1789 b	TAP2, transporter 2, ABC (ATP 1789 binding cassette)	GAACAACATT [G/A] CTTATGGGCT	ტ <u>გ</u>	A	Æ	H
G440u5	WIAF-11819	M74447	1565 1	TAP2, transporter 2, ABC (ATP 1565 binding cassette)	AAGGGGCTGA [C/T] GTTTACCCTA	υ Σ	Ħ	H	Σ
G440u6	WIAF-11820	M74447	1254 b	TAP2, transporter 2, ABC (ATP 1254 binding cassette)	TGCACTTGGG [G/T] GTGCAGATGC	ა ე	H	Ŋ	ტ
G440u7	WIAF-11788	M74447	1231  k	TAP2, transporter 2, ABC (ATP 1231 binding cassette)	GTACCTGCTC [A/G] TAAGGAGGGT	Ψ A	ტ	н	۸
G440u8	WIAF-11821	M74447	1404 1	TAP2, transporter 2, ABC (ATP 1404 binding cassette)	TGCTCAGCAA [C/T] GTGGGAGCTG	ე დ	E	z	z

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G440u9	WIAF-11783	M74447	2187	TAP2, transporter 2, ABC (ATP 2187 binding cassette)	ccceccrear [1/6] caecaeceec	S E		Þ	Þ
G440u10	WIAF-11786	M74447	1825	TAP2, transporter 2, ABC (ATP 1825 binding cassette)	rgataaggtg [a/g] rggcggcrgc	MA	ტ	Σ	Λ
G4400ul	WIAF-14007	HT97396	839 A33	A33	GCCAATCAAA [G/T] GAGGGCTCAC	M	H	×	Z
G4404u1	WIAF-14013	HT1215	109	ACP2, acid phosphatase 2, 109 lysosomal	ccgcccaccc[g/a] ggcccggagT	ტ ∑	Æ	씸	ŏ
G4404u2	WIAF-14016	HT1215	1271	ACP2, acid phosphatase 2, lysosomal	ACCGCCACGT [C/T] GCAGATGGGG	S C	F	Þ	>
G4406u1	WIAF-13661	HT3564	872	872 ACPP, acid phosphatase, prostate	ACAAAAAACT [T/C] ATCATGTATT	S	Ü	니	니
G4406u2	WIAF-13662	HT3564	839	839 ACPP, acid phosphatase, prostate	ATCACATGAA [G/A] AGAGCAACTC	<u>ი</u>	Ø	ᅜ	×
G4406u3	WIAF-13881	HT3564	741	741 ACPP, acid phosphatase, prostate	AGAATTGTCA [G/T] AATTGTCCCT	Z U	H	[11]	*
G441u1	WIAF-10166	M77349	869	TGFBI, transforming growth factor, beta-induced, 68kD	GTGCCCGGCT [C/G] CTGAAAGCCG	ა ე	v	д	ഥ
G441u2	WIAF-10168	M77349	1028	TGFBI, transforming growth	GGCTGTCTGT [A/G] GAGACCCTGG	ง 4	Ö	>	Þ
G4 <b>41</b> u3	WIAF-10169	M77349	1667	TGFBI, transforming growth	ACACAGTCTT [T/C] GCTCCCACAA	S	C	ഥ	ĪΨ
G441u4	WIAF-10171	M77349	1463	TGFBI, transforming growth	GTAATAGCCT [C/T] TGCATTGAGA	ა ა	H	ㅂ	ᅯ
G4411u1	WIAF-14005	HT97468	492	492 acyl-CoA	GCTGACCAAT [A/G] AGGCCACCCT	M	ŋ	×	Œ
G4411u2	WIAF-14008	HT97468	1076	1076 acy1-CoA	TGCCCGAGAC [C/T] GAGGACGAGA	ည	H	H	Н
G4412u1	WIAF-13576	HT1882	657	ACADS, acyl-Coenzyme A dehydrogenase, C-2 to C-3 short 57 chain	GCAAAACAAG [G/A] GCATCAGTGC	M	Æ	ಲ	ω
G4412u2	WIAF-13579	HT1882	1022	ACADS, acyl-Coenzyme A dehydrogenase, C-2 to C-3 short 1022 chain	TGACCTGGCG [C/T] GCTGCCATGC	S C	H	区	ద
G4415u1	WIAF-14080	HT2503	2170	acyl-Coenzyme A:cholesterol 70 acyltransferase	TCATTATATT [C/T] GAGCAGATTC	S C	E		[II4
G4415u2	WIAF-14081	HT2503	1993	acyl-Coenzyme A:cholesterol	TITCAGIICC[C/I]IAITITCIGI	8	E	Д.	д

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G4415u3	WIAF-14098	HT2503	2006	acyl-Coenzyme A:cholesterol 2006 acyltransferase	TITICIGITI [C/G] AACAIIGGCG	U E	<u> </u>	Q	<u>E</u>
G4415u4	WIAF-14101	HT2503	2365 8	acyl-Coenzyme A:cholesterol 2365 acyltransferase	GGGGTTAIGT [C/I] GCTAIGAAGI	<u>ນ</u>	<u> </u>	>	>
G4417u1	WIAF-13819	HT0542	356	AOAH, acyloxyacyl hydrolase 356 (neutrophil)	TCCAGCCAAC [G/A] ATGACCAGTC	± 5	A	Д	Z
G4417u2	WIAF-13820	HT0542	340	AOAH, acyloxyacyl hydrolase 340 (neutrophil)	TTCAGTCCTC [6/A] GCCTCTCCAG	ა ნ	A	S	လ
G4417u3	WIAF-13824	HT0542	1595	AOAH, acyloxyacyl hydrolase (neutrophil)	GCTAAATAAA [G/A] ACATGACCTA	Σ		A	z
G4417u4	WIAF-13841	HT0542	382	AOAH, acyloxyacyl hydrolase 382 (neutrophil)	CCAGCCTCTC [G/A] AATGGGCACA	S G	Æ	ശ	<u>.</u>
G4417u5	WIAF-13842	HT0542	458	AOAH, acyloxyacyl hydrolase (neutrophil)	CAACTCGACG [G/A] TCCAGGCCTC	M	Æ	Λ	— Н
G4417u6	WIAF-13843	HT0542	1201	AOAH, acyloxyacyl hydrolase (neutrophil)	GATTTCTGGA [C/T] TCCACTGTTG	დ ე			
G4417u7	WIAE-13844	HT0542	1321	AOAH, acyloxyacyl hydrolase 1321 (neutrophil)	ACCTGAAGAA [A/G] TTTATAGAAA	Ω ≰	Ü		×
G4417u8	WIAF-13845	HT0542	1404	AOAH, acyloxyacyl hydrolase 1404 (neutrophil)	GATGTCTGCA [G/A] TGGGAAGAGT	Σ	<u>გ</u>	S	Z
G4417u9	WIAF-13846	HT0542	1759	AOAH, acyloxyacyl hydrolase 1759 (neutrophil)	AATTTACAAA [C/T] TTCAATCTTT	တ	H ن		z
G4417u10	WIAF-13847	HT0542	1644	AOAH, acyloxyacyl hydrolase 1644 (neutrophil)	CTCCAGGTCA [G/A] CCCCTGCCAC	MG	¥	S	Z
G442u1	WIAF-11828	M94582	933	IL8RA, interleukin 8 receptor, alpha	CACATCGACC [G/A] GGCTCTGGAT	M	G A	~ ~	Q
G442u2	WIAF-11829	M94582	721	ILBRA, interleukin 8 receptor, 721 alpha	TCATCGTGCC[A/G]CTGCTGATCA	8	A G		д
G442u3	WIAF-11780	M94582	1027	ILBRA, interleukin 8 receptor, 1027 alpha	GCCATGGACT [C/T] CTCAAGATTC	S	H ن	ㅁ	- , <del>, i</del>
G442u4	WIAF-11792	M94582	78	IL8RA, interleukin 8 receptor, alpha	ATGGAGAGTG [A/G] CAGCTTTGAA	Σ	A G	Д	හ
G4423u1	WIAF-13752	HT2216	71.	71 ADSL, adenylosuccinate lyase	GCTATGCCAG [C/T] CCGGAGATGT	ω	H U	S	ω l
G4423u2	WIAF-13794	HT2216	126	126 ADSL, adenylosuccinate lyase	ATGGCGGCAG [C/T] TGTGGCTGTG	w	C	ㅂ	H
G4423u3	WIAF-13795	HT2216	674	674 ADSL, adenylosuccinate lyase	AGCTTGACAA [G/A] ATGGTGACAG	ß	G A	×	×

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G4428ul WIAF-13954 G444ul WIAF-10172 G444ul WIAF-10181 G444ul WIAF-10181 G445ul WIAF-10183 G445ul WIAF-13629 G446ul WIAF-13629	HT97524	F-U,	-uc					
		57 x	related protein; adipophilin	TGGTCAACCT [G/A] CCCTTGGTGA	S G	Ø	IJ	П
	HT0863	551 4	or 3	TCTGGAGACA [C/T] TACTTCCAGA	ა ა	H	耳	н
	U28694	398 1	CCR3, chemokine (C-C motif) 398 receptor 3	CGAGATCTTT [T/G] TCATAATCCT	E E	ტ	Ĺτι	۸
		2,7	CCR3, chemokine (C-C motif)	TCCTCATAAA [A/G]TACAGGAGGC	S A	ტ	×	×
	U28834 HT1392	136 7		GCAAGAAGAT [A/C] CTGCTGCCCG	S	U_	н	H
	U40373	319	Human cell surface glycoprotein 319 CD44 mRNA, complete cds.	TAGAAGGGCA [C/T] GTGGTGATTC	ე ვ	T	н	н
	HT0626	796 1	ALDOC, aldolase C, fructose-796 bisphosphate	CCCTGCTCAA [G/A] CCCAACATGG	ა ე	Æ	м	×
	U64198	754 1	IL12RB2, interleukin 12 receptor, 754 beta 2	TGAAGCCTTC[C/G]CATGTAATTT	დ ე	Ü	ഗ	ß
G446u2   WIAF-11795	U64198	1L12F 2569 beta	IL12RB2, interleukin 12 receptor, beta 2	TTTTCTCAAC [G/A] CATTACTTCC	ა ზ	4	Ŀ	E
	U64198	11.12R 2500 beta	<pre>IL12RB2, interleukin 12 receptor, beta 2</pre>	TGCAAGGTAA [A/G] GCCAATTGGA	S	ტ	<b>X</b>	×
	U64198	1918	IL12RB2, interleukin 12 receptor, 1918 beta 2	CTCCTCGCCA[G/C]GTCTCTGCAA	უ ლ	υ	Ø	Ħ
	U64198	991	IL12RB2, interleukin 12 receptor, beta 2	GTGGAGCAGA [G/A] ATCTTCGTTG	გ	Æ	図	臼
	U64198	2469	IL12RB2, interleukin 12 receptor, 2469 beta 2	AGTTCCCACG[G/C]AAATGAGAGG	<u>დ</u>	υ	ש	Ø
	U64198	1964	IL12RB2, interleukin 12 receptor, 1964 beta 2	GGTGACTTGG [C/g] AGCCTCCCAG	Σ Σ	מ	_ a	四
	U64198	2060	IL12RB2, interleukin 12 receptor, 2060 beta 2	TCTAAACTGG [C/G] TACGGAGTCG	υ Σ	<u> </u>		>
G447u1 WIAF-11796	 X03663	384	CSFIR, colony stimulating factor 1 receptor, formerly McDonough feline sarcoma viral (v-fms) oncogene homolog	CCAGTGTCCC [C/T] GAGCTGGTCG	ပ တ	E+	Δı	മ
G447u2 WAF-11836	X03663	1026	CSFIR, colony stimulating factor 1 receptor, formerly McDonough feline sarcoma viral (v-fms) 1026 oncogene homolog	ACAACAACAC [T/C] AAGCTCGCAA	<u>α</u>	ַ	E-1	H

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G447u3	WIAF-11837	X03663	886	CSF1R, colony stimulating factor 1 receptor, formerly McDonough feline sarcoma viral (v-fms) 886 oncogene homolog	GCTGAAAGTG[C/A]AGAAAGTCAT M C P	4	α	×
				CSFIR, colony stimulating factor 1 receptor, formerly McDonough feline sarcoma viral (v-fms)				
G447u4	WIAF-11797	X03663	2425	2425 oncogene homolog FUCA1, fucosidase, alpha-L- 1,	GAAGAAATAT [G/A] TCCGCAGGGA M G P	4	>	н
G4473u1	WIAF-13904	HT1352	860	J	TTCAAGCCAC[A/G]GAGCTTGCCA M A G	Ŋ	ď	24
G4473u2	WIAF-13916	HT1352	440	FUCAl, fucosidase, alpha-L- 1, 440 tissue	ACAAACTGGC[C/T]GAGTCCTGTG M C T	E4	ы	ы
G4479u1	WIAF-13637	HT1995	2465	AMPD2, adenosine monophosphate 2465 deaminase 2 (isoform L)	GCCTCAAIGA[G/T]CCIGGTCCAI - G 1	E	ı	
G4479u2	WIAF-13866	HT1995	1258	AMPD2, adenosine monophosphate 1258 deaminase 2 (isoform L)	TGGATGTGCA[T/C]GCGGACAGGA S T C	ŭ	王	н
G4479u3	WIAF-13867	HT1995	1280	AMPD2, adenosine monophosphate 1280 deaminase 2 (isoform L)	CACTITCCAI[C/I]GCTITGACAA M C 1	H	ద	U
G4479u4	WIAF-13868	HT1995	1201	AMPD2, adenosine monophosphate	TGCGGGAGGT[C/T]TTTGAGAGCA S C 1	Đ	Δ	٥
G4479u5	WIAF-13869	HT1995	1579	AMPD2, adenosine monophosphate 1579 deaminase 2 (isoform L)	GTACCAAGGG[C/T]CAGCTGGCCA S C 7	E	ro O	U
G4492u1	WIAF-14084	HT3390	866	ANX11, annexin XI (56kD 866 autoantigen)	ccrededeagr[c/T]gcrccaacaa M c :	E	ద	υ
G4492u2	WIAF-14085	HT3390	850	ANX11, annexin XI (56kD 850 autoantigen)	AGGCCATCAT[T/C]GACTGCCTGG S T	υ	н	н
G450u1	WIAF-10170	X85740	1196	CCR4, chemokine (C-C motif) 1196 receptor 4	TCCAAATTTA[C/T]TCTGCTGACA S C	H	×	×
G4502u1	WIAF-13510	HT4840	165	165 ASS, argininosuccinate synthetase	synthetase AAGGCTATGA[C/T]GTCATTGCCT S C	E+	Ω	Д
G4502u2	WIAF-13511	HT4840	369	369 ASS, argininosuccinate synthetase	Synthetase GGCCCTGCAT[C/T]GCCCGCAAAC S C	- F	н	н
G4502u3	WIAF-13512	HT4840	73	73 ASS, argininosuccinate synthetase AATCCCAGAC[G/A]CTATGTCCAG	U I	_ ≮		_

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G4502u5 WI G4502u6 WI G4502u7 WI		HT4840	129 ASS,		argininosuccinate synthetase TGGACACCTC[G/C]TGCATCCTCG	rggacacctc [g/c] rgcatcctcg	S	ر و	<u>ه</u>	3
	WIAF-13514	HT4840	285	ASS, a	argininosuccinate synthetase	AGTITIGIGGA [G/A] GAGITCAICI	ß	5	A E	<u>ы</u>
	WIAF-13515	HT4840	234 ASS,		argininosuccinate synthetase	synthetase AGGCACTGAA [G/A] CTTGGGGCCA	യ	U U	A K	M
	WIAF-13516	HT4840	316 ASS		argininosuccinate synthetase ccaGTCCAGC[G/A]CACTGTATGA	CCAGTCCAGC [G/A] CACTGTATGA	Σ	 	A A	E-1
G4502u8 WI	WIAF-13537	HT4840	426 4	ASS,	argininosuccinate synthetase	synthetase TGTCCCACGG[C/T]GCCACAGGAA	ß	υ	٦ ا	ש
G4502u9 WI	WIAF-13538	HT4840	530 7	ASS,	argininosuccinate synthetase	synthetase GAATTCTACA [A/G] CCGGTTCAAG	Σ	₹	U U	ß
G4502ul0 WI	WIAF-13539	HT4840	750 ASS,		argininosuccinate synthetase	synthetase [TTCTCGAGAT [C/T] GAGTTCAAAA	လ	ر ت	H	H
G4502u11 WI	WIAF-13540	HT4840	960 ASS,		argininosuccinate synthetase	synthetase ATGCTCATT[A/G]GACATCGAGG	S	A A	G L	- I
G4508ul WI	WIAF-13663	HT28557	1767 ARSD,	ARSD,	arylsulfatase D	CAGITITICCA [I/C] GAGCAACAIC	Z	TC	E	T
G4508u2 WI	WIAF-13693	HT28557	433 2	433 ARSD,	arylsulfatase D	TTCAGTGGAA [C/T] GCAGGCTCAG	വ	ט	Z E	Z
G4508u3 WI	WIAF-13694	HT28557	747	747 ARSD,	arylsulfatase D	GGTTTCTTCT [C/G] TGTCTCCGCG	Σ	U	Ω S	0
G4508u4 WI	WIAF-13696	HT28557	1012 ARSD	ARSD,	arylsulfatase D	CCACGAGTGC [A/G] TTCCTGGGGA	ß	A	G	A
G4508u5 WI	WIAF-13697	HT28557	1302	ARSD,	arylsulfatase D	CGAGTGATTG [G/A] AGAGCCCACG			A G	EI CD
G4508u6 WI	WIAF-13698	HT28557	1285 ARSD,	ARSD,	arylsulfatase D	GGGTGCTCCC [G/A] GCCGGCCGAG	ω	G 7	A P	
G4508u7 WI	WIAF-13699	HT28557	1807	ARSD,	arylsulfatase D	AGCCGTGCTG [C/T] GGACATTTCC	ß	ט	D H	0
G4508u8 WI	WIAF-13718	HT28557	483	483 ARSD,	arylsulfatase D	GCAAGAATCT [T/C] GCAGCAGCAT	Σ	ř	C	S
7.4 K T T W	12809	HT3430	ر 1		ASPA, aspartoacylase (aminoacylase 2. Canavan disease)	ACAACACCAC [C/T] TCTAACATGG	ഗ	ט	E E	F L
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G4518u2 WI	WIAF-13810	HT3430	851		van disease)	AAGTTGATTA[C/T]CCCCGGGATG	ഗ	บ	Y.	7 7
G4518u3 WI	WIAF-13811	HT3430	787	ASPA, (amino	ASPA, aspartoacylase (aminoacylase 2, Canavan disease)	CATCATTTCA [A/G] TGAAGGAAAA	Σ	ď	ر ن	N W
G4518u4 WI	WIAF-13837	HT3430	618	ASPA, (amino	ASPA, aspartoacylase 618 (aminoacylase 2, Canavan disease)	ACCCTGCTAC [G/A] TTTATCTGAT	Σ	ro .	A	N A
G452a1 WI	WIAF-10509	HT0695	553	APOA4,	apolipoprotein A-IV	ACCCAGGTCA [A/G] CACGCAGGCC	Σ	A	ט	N N
G452a2 W	WIAF-13124	HT0695	563	563 APOA4,	apolipoprotein A-IV	ACACGCAGGC [C/T] GAGCAGCTGC	ß	υ	T.	A

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WIAF-14131 HT1541 153 cardiac muscle subunit, isoform 1, complex, alpha subunit, isoform 1, armsporting, mitochondrial F1 complex, alpha subunit, isoform 1, transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitochondrial F1 transporting, mitar-10147 HT0768 2357 factor receptor, beta polypeptide pDGFRB, platelet-derived growth witar-10151 HT0768 3508 factor receptor, beta polypeptide pDGFRB, platelet-derived growth witar-10151 HT0768 156 factor receptor, beta polypeptide pDGFRB, platelet-derived growth witar-10161 HT0768 3346 factor receptor, beta polypeptide pDGFRB, platelet-derived growth witar-10161 HT0768 3386 factor receptor, beta polypeptide pDGFRB, platelet-derived growth witar-10161 HT0768 3386 factor receptor, beta polypeptide and witar-10161 HT0768 3386 factor receptor, beta polypeptide and witar-10161 HT0768 3386 factor receptor, beta polypeptide and witar-10161 HT0768 3386 factor receptor, beta polypeptide and witar-10161 HT0768 3386 factor receptor, beta polypeptide and witar-10161 HT0768 3438 minuit D, vacuolar proton pump) witar-13869 HT3556 654 31kD	G4524u1	WIAF-14120	HT1541	726	ATP synthase, H+ ting, mitochondrial F1 alpha subunit, isoform muscle		Σ A	<u>6</u>	н	>
HT4994	G4524u2	WIAF-14131	HT1541	153	ATP sy ting, m alpha muscle		υ Σ	H	Æ	S
PDGFRB, platelet-derived growth	G4526u1	WIAF-14130	HT4994	400	ATP synthase, H+ sing, mitochondrial delta subunit		H H	ū	>	Æ
PDGFRB, platelet-derived growth	G453u1	WIAF-10138	HT0768	1747		CTGCCGCCCA [C/T] GCTGCTGGGG	υ Σ	H	E	Σ
PDGFRB, platelet-derived growth	G453u2	WIAF-10147	HT0768	2957	platelet receptor,	TTTTGCCTTT [A/G] AAGTGGATGG		r.	H	ᆈ
WIAF-10149 HT0768 457 factor receptor, beta polypeptide WIAF-10151 HT0768 1505 factor receptor, beta polypeptide WIAF-10153 HT0768 3446 factor receptor, beta polypeptide WIAF-10161 HT0768 2030 factor receptor, beta polypeptide ATP synthase, H+ transporting, ATP synthase, H+ transporting, ATP6E, ATPase, H+ transporting, 11 WIAF-13569 HT3556 654 31kD	G453u3	WIAF-10148	HT0768	3608	PDGFRB, platelet-derived growth factor receptor, beta polypeptide	AGCCGGAGCC [A/G] GAGCTGGAAC	۵ لا	უ	Д	Д
WIAF-10151	G453u4	WIAF-10149	HT0768	457	DDGFRB, platelet-derived growth factor receptor, beta polypeptide	CAGGGCCTGG [T/G] CGTCACACCC	Σ	ro O	>	ტ
PDGFRB, platelet-derived growth	G453u5	WIAF-10151	HT0768	1505	PDGFRB, platelet-derived growth factor receptor, beta polypeptide	AGCTGACACT [G/C] GTTCGCGTGA	ა ზ	Ü		<u> </u>
PDGFRB, platelet-derived growth   WIAF-10161   HT0768   2030 factor receptor, beta polypeptide   ATP synthase, H+ transporting,   ATP synthase, H+ transporting,   ATP synthase, H+ transporting,   ATP6E, ATPase, H+ transporting,   Iysosomal (vacuolar proton pump)   WIAF-13569   HT3556   654 31kD	G453u6	WIAF-10153	HT0768	3446	platelet receptor,	ACCCCAAACC[C/T]GAGGTTGCTG	_ນ	E	<u> </u>	Д,
ATP synthase, H+ transporting,   MIAF-13616   HT1618   343 subunit D, vacuolar   ATP6E, ATPase, H+ transporting,   Iysosomal (vacuolar proton pump)   WIAF-13569   HT3556   654 31kD	G453u7	WIAF-10161	HT0768	2030	platelet receptor,	TTGGCAGAA [G/A] AAGCCACGIT	დ დ	A.	×	ೱ
ATP6E, ATPase, H+ transporting, lysosomal (vacuolar proton pump) WIAF-13569 HT3556 654 31kD	G4533u1	WIAF-13616	HT1618	343	ATP synthase, H+ subunit D, vacuo]	GTTACATGAT [C/T] GACAACGTGA	ა ე	H		
	G4534u1	WIAF-13569	HT3556	654	E,	TAAAGGTTTC [C/T] AACACCCTGG	ى د	E-1	ω.	ω

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G4535u1	WIAF-13747	HT27972	357	ATP50, ATP synthase, H+ transporting, mitochondrial F1 complex, 0 subunit (oligomycin 357 sensitivity conferring protein)	TCACTACCAA [C/T] CTGATCAATT	ى ت	<u> </u>	Z	z
G4535u2	WIAF-13748	HT27972	144 8	ATP50, ATP synthase, H+ transporting, mitochondrial F1 complex, O subunit (oligomycin sensitivity conferring protein)	aggtatacgg [T/C] attgaaggtc	N H	υ	ප	യ
G4535u3	WIAF-13792	HT27972	329 s	ATP50, ATP synthase, H+ transporting, mitochondrial F1 complex, O subunit (oligomycin sensitivity conferring protein)	atcacagcaa [a/g] agagaggttc	м д		M	꿈
G4539u1	WIAF-13711	HT48520	288	288 ATPase, 14 kDa subunit, vacuolar	TGCCCTGGAC [G/A] CCCACCAGCA	M	Ą	A	H
G4548u1	WIAF-14127	HT1574	3138 1	ATPase, Ca2+ transporting, plasma 3138 membrane, isoform 2	CGCAATGTCT [T/C] TGACGGCATC	M	ບ	Щ	κĵ
G4548u2	WIAF-14137	HT1574	2089	ATPase, Ca2+ transporting, plasma 2089 membrane, isoform 2	GCACTATCTG [C/T] GTGGCCTACC	ສ ແ	Ę÷	Ü	บ
G4548u3	WIAF-14140	HT1574	2924	ATPase, Ca2+ transporting, plasma 2924 membrane, isoform 2	CAGGACCATG [A/T] TGAAGAACAT	Σ	E	_ Σ	н
G4549u1	WIAF-14161	HT1346	524	ATP2B4, ATPase, Ca++ 524 transporting, plasma membrane 4	TGCACTGACC[C/T]AGATTAATGT	N D	E	α	*
G4549u2	WIAF-14162	HT1346	715	ATP2B4, ATPase, Ca++ 715 transporting, plasma membrane 4	ATGTCACGCT [C/T] ATCATCCTGG	S S	€-1		Ц
G4549u3	WIAF-14163	HT1346	208	ATP2B4, ATPase, Ca++ 508 transporting, plasma membrane 4	AGCTGCGTTC[G/A]AGGGATGCAC	დ ტ	A	S	ß
G4549u4	WIAF-14166	HT1346	1084	ATP2B4, ATPase, Ca++ 1084 transporting, plasma membrane 4	TGATCCAAGG [G/A] AATGATCTGA	ν σ	Æ	ט	₀
G4552u1	WIAF-13630	HT0867	710	ATP7A, ATPase, Cu++ transporting, alpha polypeptide (Menkes 710 syndrome)	TACTAGCACT [A/G] TTGAAGGAAA	M	<u></u>	<u> </u>	Δ

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G456u2	-MTAK-100/4								
G456UZ	1000		408 EDNI,	endothelin i				7 5	7 [
	WIAF-10075	34	PRP EDNI,	endothelin l	CAGACCG1GA [A/G] AATAGATGCC			1	7
G456a3	WIAF-10507	HT2834 8	861 EDN1,	endothelin 1	TGAAAGGCAA [T/G] CCCTCCAGAG	E	Ü	×	z
G4565u1	WIAF-14041	HT28561	ATE 320 tre	ATP1G1, ATPase, Na+/K+ 320 transporting, gamma 1 polypeptide	CGAGGCTGCT [G/A] TTACGGCTCA	S	A	<u> </u>	<u> </u>
G4565u2	WIAF-14062	HT28561 2	ATI 216 tra		CAGTGACGGG [G/A] ACAAAGGTCT	<u>ب</u> ق	4	Α	Z
G4565u3	WIAF-14063	HT28561	ATE 315 tra	ATP1G1, ATPase, Na+/K+ transporting, gamma 1 polypeptide	ACCGCCGAGG [C/A] TGCTGTTACG	ບ <b>∑</b>	A	<u>_</u>	Σ
G4565u4	WIAF-14064	HT28561	ATE 531 tra	ATP1G1, ATPase, Na+/K+ transporting, gamma 1 polypeptide	TTTCCCCAGG[T/C]GAATGGGCTG	N	υ .	*	<u>r</u>
G4568u1	WIAF-14212	HT0082	AMFR, a	utocrine motility factor	TGCCTCATGC [A/G] TACGTCCCAC	M	Ü	Н	Δ
G457a1	WIAF-10489	HT2903	SELL, 321 adhes	selectin L (lymphocyte	ACAAATCTCT [C/T] ACTGAAGAAG	8 D	E-1	ᄀ	그
G457a2	WIAF-10490	HT2903	SEI 577 adì	SELL, selectin L (lymphocyte adhesion molecule 1)	CCAGIGICAG [T/C] TIGIGAITCA	E X	<u> </u>	[I4	ㅁ
G457a3	WIAF-10491	HT2903	SEI 601 adł	SELL, selectin L (lymphocyte adhesion molecule 1)	TGAGCCTTTG [G/C] AGGCCCCAGA	M	U ZD	田	Ø
G457a4	WIAF-10492	HT2903	SEI 637 adł	SELL, selectin L (lymphocyte adhesion molecule 1)	CTGTACTCAC[C/T]CTTTGGGAAA	Σ	F	Ъ	လ
G4573u1	WIAF-13568	HT28320	MG2 913 943 ace	MGAT2, mannosyl (alpha-1,6-)- glycoprotein beta-1,2-N- acetylglucosaminyltransferase	CGGACAACCT[G/T]ACGCTGCGGT	رن د	E+1	卢	<u> </u>
G4574u1	WIAF-13805	HT0198		beta-1,4 N- 63 acetylgalactosaminyltransferase	CGGCCTCCGG [C/G] TACCTCTTGC	Σ	<u>ဗ</u>	니	>
G4574u2	WIAF-13806	HT0198	be: 415 ac	beta-1,4 N- 415 acetylgalactosaminyltransferase	TGCCACAAGA [G/A] AGCAGGAGTT	υ Σ	A	<u> </u>	×

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G4574u3	WIAF-13807	HT0198	726	beta-1,4 N-726 acetylgalactosaminyltransferase	AACTACAACT [G/T] GTCACTTACA	D E	<u>, , , , , , , , , , , , , , , , , , , </u>	
G4574u4	WIAF-13836	HT0198	559	beta-1,4 N- 59 acetylgalactosaminyltransferase	AGGGCTGAGC [C/A] TTCAGGCAGC M	D A	H	H
G4575u1	WIAF-13626	HT0341	1251	GCNT1, glucosaminyl (N-acetyl) transferase 1, core 2 (beta-1,6-N-acetylglucosaminyltransferase)	AGTATGATCT [A/G] TCTGACATGC	\ \ \	ы	Ц
G4577u1	WIAF-13971	HT1495	1268	SIAT1, sialyltransferase 1 (beta-galactoside alpha-2,6-sialytransferase)	ATTTCTTTAA [C/T] AACTACAAGA	H ن	Z	
G458u1	WIAF-10063	HT2968	1464 ALB,	ALB, albumin	GTGCAGAAGA [C/A] TATCTATCCG M	C		Œ
G458u2	WIAF-10089	HT2968	1470 ALB,	ALB, albumin	AAGACTATCT [A/C] TCCGTGGTCC S	A C	Ä	ᆈ
G458u3	WIAF-10091	HT2968	1707 ALB,	ALB, albumin		C	-	ы
G458a4	WIAF-10504	HT2968	889	889 ALB, albumin				[IL4
G458a5	WIAF-10508	HT2968	1475 ALB,	ALB, albumin				Œ
G458a6	WIAF-12091	HT2968	1330	1330 ALB, albumin		r U		
G458a7	WIAF-12092	HT2968	1408	1408 ALB, albumin	CCTAGGAAAA [G/a] TGGGCAGCAA M	ro Co	a	Σ
				branched-chain keto acid				
74592111	WTAF-14126	HT2128	985	dehydrogenase E1, alpha polvpeptide	ACCAGCCCTT [T/C] CTCATCGAGG	E	C C	Ex.
3				BARD1, BRCA1 associated RING				
G4593ul	WIAF-13574	HT97373	1743	1743 domain 1	GCTAGCCACT [G/C] CTCAGTAATG M	ซ	ت ت	တ
G4593u2	WIAF-13592	HT97373	1167	BARD1, BRCA1 associated RING	TGTTCTTCAC[C/T]ACCTTCATGC M	υ	T.	
G4593u3	WIAF-13593	HT97373	1591	BARD1, BRCA1 associated RING domain 1	AGAATGGGCA[C/T]GTGGATATAG	ບ	H	Ħ
G4593u4	WIAF-13594	HT97373	2030	BARD1, BRCA1 associated RING 2030 domain 1	AAAGTATGAA [A/G] TTCCTGAAGG	A	G	>
24503115	WT 2 F - 1 3 F 9 F	HT97373	2006	BARD1, BRCA1 associated RING	AAGAAAAGTA [T/C] GTGAACAGGA	E	ن ن	ద
1.100 P. P. P. P. P. P. P. P. P. P. P. P. P.	WTAF-13920	HT4273	1803	CDH13, (heart)	TCGTACCCGA [C/T] GTCTCCTACG	ŭ	O F	Ω
G461411	WIAF-13733	HT4835	91	S100A3, S100 calcium-binding protein A3	AGGATGGCCA [G/A] GCCTCTGGAG M	r _D	A	×
G4614u2	WIAF-13734	HT4835	203	S100A3, S100 calcium-binding protein A3	TGCTGCAGAA [G/A] GAGCTGGCCA	ט	A K	M
G4614u3	WIAF-13769	HT4835	344	S100A3, S100 calcium-binding	TCTACTGCCA [C/T] GAGTACTTCA	ŭ	H	<u>н</u>

G462n1	WIAF-10134	HT4753	600 t	PDGFA, platelet-derived growth	ACGGGGTCCA[C/T]GCCACTAAGC	ט	EH	H	н
G4627u1	WIAF-14042	HT0771	186 2	annexin VI (p68)	GGAGGCCATA [C/T] TGGACATAAT	ບ	E	П	ы
G4627u2	WIAF-14043	HT0771	1664 ANX6,	annexin VI (p68)	CAGACACACC [T/C] AGTGGAGACA S	E	ပ	Д	д
G4627u3	WIAF-14067	HT0771	1498 7	ANX6, annexin VI (p68)	AAGGAGGACT [A/G] TCACAAGTCC M	A.	υ	×	ŭ
G4644ul	WIAF-13801	HT1736	1990	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	TGGTGGAGAA [G/A] TCAGTGACAG	ro l	Ą	X	M
G4644u2	WIAF-13802	HT1736	1866	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	ATTGGCTACC[C/T]AGTGATGATC M	<u></u> υ	Ð	д	П
G4644u3	WIAF-13803	HT1736	1993	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	TGGAGAAGTC [A/C] GTGACAGGTT	4	U	S	Ω
G4644u4	WIAF-13804	HT1736	1860	CPS1, carbamoyl-phosphate 1860 synthetase 1, mitochondrial	GACACCATTG [G/A] CTACCCAGTG M	Ď	Æ	ro l	Д
G4644u5	WIAF-13831	HT1736	1087	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	AGCCTGTTTT [G/T] AATATCACAA	D F	EH	н	Ē4
G4644u6	WIAF-13835	HT1736	1958	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	CACAAAGGCC[T/C]TTGCTATGAC M	H	υ	(Z1 ₄	L L
24644117	™ 7477 7277	HT1736	1332	CPS1, carbamoyl-phosphate synthetase 1, mitochondrial	M AAAGCTACCA [C/A] CATTACATCA	ט	4	E	'n
G4659u1	WIAF-14143	HT1183	1830	catenin, alph	GTGCCAACGT[T/C]CCTCAACCGT S	H S	บ	>	>
G466u1	WIAF-10164	896000	2403	SREBF1, sterol regulatory element binding transcription factor 1	AGCAGTGCCC [G/A] CCAGGCCTGC M	ک ن	< 4		Ħ
G4662u1	WIAF-13710	HT2142	2183	CTNNB1, catenin (cadherin- 2183 associated protein), beta 1 (88kD)	(88kD) TITIGITCCG[A/C]AIGICIGAGG S	ر لا	บ	ĸ	P¥.
G467a1	WIAF-13304	X72861	827	ADRB3, adrenergic, beta-3-, 827 receptor	GGCCATCGCC [T/C] GGACTCCGAG	E E	U	<b>Z</b>	ద
G467a2	WIAF-13305	X72861	832	ADRB3, adrenergic, beta-3-, receptor	TCGCCTGGAC[T/A]CCGAGACTCC	E⊣ (S)	Æ	E⊣	E
G467a3	WIAF-13306	X72861	870	ADRB3, adrenergic, beta-3-, 870 receptor	TTCGTGACTT [C/T] GCTGGCCGCA	υ Σ	E+	Ω	П
G467a4	WIAF-13307	X72861	1761	ADRB3, adrenergic, beta-3-, receptor	TGCGCCGCCG [C/I] CCGCCCGGCC	υ Σ	E⊢	4	⊳

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G467a5	WIAF-13308	X72861	1899	ADRB3, adrenergic, beta-3-, 1899 receptor	TCTGTTGATC [A/C] GAACCTGTGG	- A	೮	1	
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G46/1U1	WLAF - 13936	076TIU	TOT	(010)			) 1	-	
G4673ul	WIAF-13889	HT0191	1349	1349 CDC25A, cell division cycle 25A	TCTGGGGCCA [G/C] CCCCAAAGAG	ב ב	را	ω	-
G4674u1	WIAF-13821	HT1393	261	261 CDC25B, cell division cycle 25B	ACGACCTCGC [C/T] GGGCTCGGCA	S	_=	4	А
G4674u2	WIAF-13822	HT1393	1297	1297 CDC25B, cell division cycle 25B	GATGGTGGCC [C/T] TATTGACGGG	S	₽	ㅁ	ы
G4674u3	WIAF-13823	HT1393	1083	1083 CDC25B, cell division cycle 25B	ATAAGCGGAG [G/A] CGGAGCGTGA	S D	4	ద	ద
G4674u4	WIAF-13827	HT1393	1446	1446 CDC25B, cell division cycle 25B	AGAGCCCCAT [C/T] GCGCCCTGTA	S	E	н	н
G468a1	WIAF-13309	1.37019	192	ASIP, agouti (mouse)-signaling protein	AAATCCAAAC[C/A]GATCGGCAGA	υ Σ	4	д	Ø
G4691u1	WIAF-13753	HT97602	179	CMKBR9, chemokine (C-C motif) receptor 9	TATAGCCTGA [T/A] TTTTGTGTTG	E E	₹		z
G4691u2	WIAE-13754	HT97602	134	CMKBR9, chemokine (C-C motif) receptor 9	AAGGATGCAG [T/C] GGTGTCCTTT	M	υ	>	Æ
G4691u3	WIAF-13755	HT97602	193	CMKBR9, chemokine (C-C motif) receptor 9	TGTGTTGGGC[C/T]TCAGCGGGAA	Z U	H	ㅂ	Ēų
G4691u4	WIAF-13756	HT97602	770	CMKBR9, chemokine (C-C motif) 770 receptor 9	AAAATAGCTG [C/T] AGCCTTGGTG	МС	H	Ą	٥
G4691u5	WIAF-13759	HT97602	1130	CMKBR9, chemokine (C-C motif)	TCTGAGAACT [A/C] CCCTAACAAG	M	Ü		ß
G4691u6	WIAF-13796	HT97602	482	CMKBR9, chemokine (C-C motif) 482 receptor 9	AGGCTGAGGA [C/A] CCGGGCCAAG	Σ	A		z
G4691u7	WIAF-13797	HT97602	259	CWKBR9, chemokine (C-C motif) receptor 9	GATGGTTGAG [A/G] TCTATCTGCT	M	ರ	н	>
G4691u8	WIAF-13798	HT97602	434	CMKBR9, chemokine (C-C motif) receptor 9	ATGAGCCTGG [A/G] CAAGTACCTG	M M	ט	Д	ტ
G4691u9	WIAF-13799	HT97602	755	CMKBR9, chemokine (C-C motif) receptor 9	CAGGCCCGGG [C/T] TTTAAAAATA	Σ	H	A	>
G4699u1	WIAF-14040	HT4277	1426	BAAT, bile acid Coenzyme A: amino acid N-acyltransferase (glycine N- 1426 choloyltransferase)	TTCCAGATGT [G/T] ACCAGTCAAC	<u>ა</u>	E+	>	>

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G4726u1	WIAF-14128	HT48614	1606	AOC3, amine oxidase, copper containing 3 (vascular adhesion 1606 protein 1)	TCCACCCCAG [T/C] GGGGCCATAG	ω H	υ	S	S
G4726u2	WIAR-14129	HT48614	2242	AOC3, amine oxidase, copper containing 3 (vascular adhesion protein 1)	TTCCTAACAC [A/G] GTGACTGTGG	δ 4	υ	E	F
G4726u3	WIAF-14141	HT48614	659	AOC3, amine oxidase, copper containing 3 (vascular adhesion 659 protein 1)	CCTGCCCTAT [C/T] ACCGACGCCC	υ Σ	EH	д	Y
G4744u1	WIAF-13683	HT2599	564	CTH, cystathionase (cystathionine 564 gamma-lyase)	ATATTGTCCA[T/C]AAGCATGGAG	N E4	υ	Ħ	н
G4748u1	WIAF-14144	HT1061	242	CYBA, cytochrome b-245, alpha 242 polypeptide	GGGACAGAAG [C/T] ACATGACCGC	υ Σ	Ę	Щ	þч
G4748u2	WIAF-14145	HT1061	265	CYBA, cytochrome b-245, alpha 265 polypeptide	TGGTGAAGCT [G/C] TTCGGGCCCT	<u>თ</u>	υ U		ıл
G4750ul	WIAF-14116	HT48417	156	56 CYB5, cytochrome b-5	TGAAGTACTA[C/T]ACCCTAGAGG	S C	E-	×	×
6475111	WIAF-13770	HT1285	495	UQCRC2, ubiquinol-cytochrome c	AGAATTTCGT [C/A] GTTGGGAAGT	υ Έ	ď.	ద	ß
G4788u1	WIAF-13931	HT28249	1864 DSC3,	DSC3, desmocollin 3	CTGTTGATCC [T/C]GATGAACCTG	S E	บ	Д	Дı
G4788u2	WIAF-13933	HT28249	2000	2000 DSC3, desmocollin 3	TGGATTTCAA [G/T] AATATACCAT		£4	Œ	*
G4788u3	WIAF-13945	HT28249	2524	2524 DSC3, desmocollin 3	ACACTTACTC [G/A] GAGTGGCACA	S G	4	S	ß
G479u1	WIAF-12567	U36310	894	GPD2, glycerol-3-phosphate 894 dehydrogenase 2 (mitochondrial)	GGGAAAGTGC [A/G] TGTGAGCGGC	M M	r D	E	pt.
G479u2	WIAF-12574	U36310	1657	GPD2, glycerol-3-phosphate 1657 dehydrogenase 2 (mitochondrial)	CTGGCAAAAG [G/T] TGGCCTATTG	<u>ნ</u>	<u></u>	<u>~</u>	ß
G479u3	WIAF-12575	U36310	1131	GPD2, glycerol-3-phosphate	GTTALTITICT [1/C] CTTACCCIGG	E-I	ט	Et.	S
G480u1	WIAF-12175	HT336	250	GRB2, growth factor receptor- 250 bound protein 2	AATGAAACCA[C/A]ATCCGTGGTT	υ Σ	A	н	z
G4819u1	WIAF-13985	HT97576	1804	EYA1, eyes absent (Drosophila) 1804 homolog 1	CCCTGCACCA[T/C]GCCTTGGAAC	ω EH	υ	田	н
G482u1	WIAF-12181	J04501	1186	GYS1, glycogen synthase 1	CTGACGTCTT [T/C] CTGGAGGCAT	S	Ü	Ex.	<u> </u>

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G482112	WIAF-12195	J04501	1406	GYS1, glycogen synthase 1	CCTTCCCGAC [A/G] TGAACAAGAT	M	ტ_	Σ	Þ
G4827u1	WIAF-14177	HT97477	9 89	elongation	CGAGCTGGCC [A/G] TGATGGTGAT	M	υ	Ħ	В
G483a1	WIAF-12113	HT4341	1850 GSY2	\$\$Y2	TTACCAGCAT [G/T] CCAGACACCT	ĭ E	Н	Æ	ß
G483u2	WIAF-12148	HT4341	1130 GSY2	\$SY2	GTTTTTCATT [A/C] TGCCTGCCAA	M	ပ	Σ	ı
G483u3	WIAF-12149	HT4341	880 GSY2	SSY2	GCTTGAATGT [T/G] AAGAAATTTT	S	Ŋ	>	٥
G483u4	WIAF-12150	HT4341	1115 GSY2	SSY2	CATCACAGTG [G/A] TGGTGTTTTT	r E	Æ	>	Σ
G483u5	WIAF-12156	HT4341	1230 GSY2	SSY2	GAAAAGTTTG [G/A] AAAAAAACTC	Σ Ω	ď	ט	ы
G483u6	WIAF-12159	HT4341	2033 GSY2	SSY2	TGAGAGATAC [G/A] ATGAGGAAGA	Σ D		Д	z
G483u7	WIAF-12160	HT4341	1836 GSY2	3SY2	TACTTAGGCA [G/C] ATATTACCAG	Σ.	U	ద	ы
G483u8	WIAF-12161	HT4341	1678 GSY2	3SY2	CTTACGGTAT [T/C] TACATCGTTG	S	ပ	н	н
G483u9	WIAF-12177	HT4341	790 GSY2	35.42	GCGCTCACGT [G/C] TTCACCACGG	S D	υ	>	۸
G483u10	WIAF-12188	HT4341	1728 GSY2	SSY2	TGCAATCAGC [T/C] GACTAAGTTT	E.	ပ	크	ч
G484u1	WIAF-12151	HT5111	487 GSY3	\$SY3	CATCAAAGTG [A/G] TTGGCAATGG	M	೮	н	Þ
G484u2	WIAF-12187	HT5111	1141 GSY3	SSY3	AACCCGGGAA [C/T] AAATCCGAGA	N N	[-1	ď	*
G489u1	WIAF-12152	HT2607	1181	IRS1, insulin receptor substrate	AAGAAGTGGC [G/A] GCACAAGTCG	ზ	Ą	ద	ø
G489u2	WIAF-12184	HT2607	1031	IRS1, insulin receptor substrate	ATGGCGAGCC[C/T]TCCGGAGAGC	M	H	凸	ы
G492a1	WIAF-13345	L08603	307	307 MC4R, melanocortin 4 receptor	AGAAACCATT [A/G] TCATCACCCT	M	_ დ	<u> </u>	Δ
7. 64 9.20 7.1.2.9	WTAR-12154	X67594	346	MCIR, melanocortin 1 receptor (alpha melanocyte stimulating 346 hormone receptor)	CGCGCTGGTG [G/T] TGGCCACCAT	Z E	H	>	<u></u> <u></u>
G493u2	WIAF-12167	X67594	646	646 hormone receptor)	GACCCTGCCG [C/T] GGGCGCGGCA	Σ	E	ద	M
G493u3	WIAF-12170	X67594	1110	MC1R, melanocortin 1 receptor (alpha melanocyte stimulating 1110 hormone receptor)	AGGTGCTGAC [A/G] TGCTCCTGGT	ω 4	<b></b>	E	E
G493u4	WIAF-12186	X67594	442	MCIR, melanocortin 1 receptor (alpha melanocyte stimulating 442 hormone receptor)	CGGGAGCAAC [G/T] TGCTGGAGAC	M	E⊣	>	
G498u1	WIAF-11809	J04127	1305	CYP19, cytochrome P450, subfamily 1305 XIX (aromatization of androgens)	CTTATAGGTA[C/T]TTTCAGCCAT	ა ა	E	<u>&gt;</u>	**

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G498u2	WIAF-11810	J04127	1377	CYP19, XIX (aro	CYP19, cytochrome P450, subfamily 1377 XIX (aromatization of androgens)	TGAAAGCCAT [C/T] CTCGTTACAC		ر د	E-	H	H
G498u3	WIAF-11811	J04127	CYP1	9, (arc	9, cytochrome P450, subfamily (aromatization of androgens)	CGATTCCACG[T/C]GAAGACATTG		E E	ט	Δ	4
G498u4	WIAF-11838	J04127	1055	CYP19, XIX (aro	CYP19, cytochrome P450, subfamily 1055 XIX (aromatization of androgens)	attggtgaga [g/a] agacataaag		Σ	<u>م</u>	ద	×
7498115	WIAF-11800	J04127	1001	CYP19, XIX (aro	9, cytochrome P450, subfamily (aromatization of androgens)	ATTGCAAAGC [A/G] CCCTAATGTT		Σ	A G		ద
G4 9911	WIAF-11785	HT1439	2142	2142 ESR1, e	estrogen receptor 1	TCCCTGCCAC [A/G] GTCTGAGAGC		S	A G		E
G499u2	WIAF-11801	HT1439	443	443 ESR1, e	estrogen receptor 1	CCCCTGAACC [G/A] TCCGCAGCTC					
G500u1	WIAF-11803	X99101	793	ESR1,	estrogen receptor 1	CATGATCAGC [T/C] GGGCCAAGAA		$\neg$	T		
G500u2	WIAF-11816	X99101	489	489 ESR1, e	estrogen receptor 1	GGAAGTGTTA [C/T] GAAGTGGGAA				T	
G500u3	WIAF-11817	X99101	474	474 ESR1, e	estrogen receptor 1	AGGCCTGCCG [A/G] CTTCGGAAGT					T
G505u1	WIAF-11824	HT1113	1063	1063 PRLR, F	prolactin receptor	GCTTTGAAGG [G/A] CTATAGCATG					
G505u2	WIAF-11827	HT1113	2083	2083 PRLR, I	prolactin receptor	GCAACATCAA [G/A] CAAGTGCAGG					
G505u3	WIAF-11787	HT1113	582	582 PRLR, I	prolactin receptor	GAGGACATAC [A/G] TCATGATGGT					П
G505u4	WIAF-11802	HT1113	792		prolactin receptor	CCTGTATGAA[A/C]TTCGATTAAA		Σ	A C		H
				SRD5A1, streductase,	, steroid-5-alpha- ase, alpha polypeptide 1 (3. alpha-steroid delta 4-						
G509u1	WIAF-11789	M32313	378	dehydı	genase alpha 1)	CACTGTTGGC [A/G] TGTACAATGG	ACAATGG	တ	A	G	A.
	4 th	717 7000	n c	STAR,	STAR, steroidogenic acute	CCAATGTCAA [G/A]GAGATCAAGG	ATCAAGG	ß		¥   X	<u> </u>
GOLUGIL	WIAE-1024	HT0488	1139	inhihin	1139 inhibin, beta B	CCAACATGAT [T/C] GTGGAGGAGT	GAGGAGT	Ω	E	CI	Н
60.00 T	WTAR-13507	n31770	517	ACVR2, II	activin A receptor, type	CTTATTTTCC [G/A] GAGATGGAAG	ATGGAAG	ß	ט	A D	<u>-</u>
G520u2	WIAF-13532	D31770	AC 1177 II	ACVR2, II	activin A receptor, type	CAGCTTGCAT [T/G] GCTGACTTTG	GACTTTG	Σ	H	<u>н</u>	Σ
G520113	WIAF-13533	D31770	1189	ACVR2, II	activin A receptor, type	CTGACTTTGG [G/C] TTGGCCTTAA	GCCTTAA	S	r U	ט	ט
2520114	WTAF-13534	D31770	AC 1024 II	ACVR2, II	activin A receptor, type	TCTCTTGGAA [T/C] GAACTGTGTC	ACTGTGTC	လ	F	C N	
G523u1	WIAF-12155	HT4996	538	TR,	oxytocin receptor	TGAGCGGGAA [C/T] GCGTGTGTGC	STGTGTGC	വ	υ υ	Z E	Z

0.00	10100	TT 096	1057 OXTR	oxvtocin receptor	TCTGGCAGAA [C/T] TTGCGGCTCA	S C	H	Z	z
692342	MIRE - 12100	1,05144	190	phosphoenolpyruvate		ე	Æ	니	니
1017 COUNT	100 T T T T T T T T T T T T T T T T T T	V00572	00	PGK1. phosphoglycerate kinase 1	AAGCCACTGT [G/C] GCTTCTGGCA	හ හ	<u>U</u>	Λ	Δ
G520UI	WTAF-10307	HT0508	723	epair protein XRCC1		S	Æ	Д	д
G53u2	WIAF-10308	HT0508	746 DNA	repair protein XRCC1		υ Σ		4	⊳
G53u3	WIAF-10309	HT0508	1884	7			EH	S	လ
G53u4	WIAF-10362	HT0508	425 DNA	repair protein XRCC1				ద	Ξ
G534a1	WIAF-13310	U28281	1284	1284 SCTR, secretin receptor				z	z
G534a2	WIAF-13311	U28281	1404	1404 SCTR, secretin receptor	AGCAGAGCCA [G/A] GGCACCTGCA			의	α
G535u1	WIAF-12157	HT5001	1158 SHC1		ATGCTCTTCG [G/C] GTGCCTCCAC				ద
G535u2	WIAF-12196	HT5001	774	774 SHC1	ATGAGGAGGA [G/A] GAAGAGCCAC	დ ტ	Æ	臼	[2]
				SLC2A4, solute carrier family 2 (facilitated glucose transporter),					
G536u1	WIAF-13923	M20747	535	535 member 4	GCCTGGCCAA [C/T] GCTGCTGCCT	<u>အ</u>	=	4	4
G538u1	WIAF-11812	M55531	438	SLC2A5, solute carrier family 2 (facilitated glucose transporter), member 5	GCAGCAGAGT [C/T] GCCACATCAT	ى د	<u> </u>	> >	>
G538u2	WIAF-11813	M55531	124	SLC2A5, solute carrier family 2 (facilitated glucose transporter),	GACGCTTGTG [C/T] TTGCCCTGGC	υ Σ	F	긔	[ <del>-</del>
				SLC2A5, solute carrier family 2 (facilitated glucose transporter),					
G538u3	WIAF-11791	M55531	816		ACAGGGAGGT [G/A] GCCGAGATCC	S	<b>A</b>	>	>
G539u1	WIAF-12158	K03195	224	Human (HepG2) glucose transporter gene mRNA, complete cds.	TCATGCTGGC[T/C]GTGGGAGGAG	. E-I	U	A	4
C L	# F P P P P P P P P P P P P P P P P P P	X02195	1244	Human (HepG2) glucose transporter gene mRNA, complete cds.	CCATCGCGCT [A/G] GCACTGCTGG	S	ج 2	- I	귀
G559U2	WIAF-12134	HT960	1100	SOS1	AGTGAAGATC[A/C]AGAAGACAAG	M	A C		
G540u2	WIAF-12165	HT960	933	933 SOS1	ATGATCGTTT[C/T]CTTAGTCAGT				
G540u3	WIAF-12178	HT960	399	399 SOS1	TAGTAGCAGT[C/T]TTAGAATACA		$\neg$		
G540u4	WIAF-12193	HT960	195	195 SOS1	CTCAGCCCCG [A/C] AGTGCTTCAG				
G540u5	WIAF-12197	HT960	1329	1329 SOS1	GTTGTAATGA[A/G]TTTATAATGG				
G540u6	WIAF-12198	HT960	1339	SOS1	ATTTATAATG [G/A] AAGGAACTCT				
G543a1	WIAF-13312	300306	1373	SST, somatostatin	AAGCAGGAAC [T/C] GGCCAAGTAC	Σ	<u>၂</u>	-1	24

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G544u1	WLAF'-121/4	H12/489	700	(Hyperimsurinemia) TKT, transketolase (Wernicke-				
G546u1	WIAF-13618	HT225	426	426 Korsakoii syndrome)		,		
G551u1	WIAF-11709	HT1118	257	TNFRSF1B, tumor necrosis factor 257 receptor superfamily, member 1B	GCTGCAGCAA [A/G] TGCTCGCCGG	A	D D	×
G551u2	WIAF-11710	HT1118	449	TNFRSF1B, tumor necrosis factor 449 receptor superfamily, member 1B	TCTGCACCTG [C/T] AGGCCCGGCT S	۵	D L	υ
G551u3	WIAF-11719	HT1118	648	TNFRSF1B, tumor necrosis factor receptor superfamily, member 1B	GATCTGTAAC [G/A] TGGTGGCCAT M	ט	A A	Σ
G551u4	WIAF-11673	HT1118	676	TNFRSF1B, tumor necrosis factor 676 receptor superfamily, member 1B	AATGCAAGCA [T/G] GGATGCAGTC M	E+	Σ 	ద
25 25 25	WIRE-11720	HT1118	808	TNFRSF1B, tumor necrosis factor 808 receptor superfamily, member 1B	CCAAGCACCT [C/T] CTTCCTGCTC M	υ •	E E	[īt4
G552u1	WIAF-12229	HT5108	384	TRAP3	GCCGCTGCCC [G/A] CTCATGCTGA S	G	A	<u>Di</u>
G555u1	WIAF-12211	U94592	478	UCP2, uncoupling protein 2 478 (mitochondrial, proton carrier)	CGCGCTACAG [T/C] CAGCGCCCAG	H	ر د	A
G556u1	WIAF-11804	AF001787	480	UCP2, uncoupling protein 2 480 (mitochondrial, proton carrier)	TCGGCCTCTA [T/C] GACTCCGTCA	EH SS	C	<u> </u>
G556u2	WIAF-11805	AF001787	563	UCP2, uncoupling protein 2 563 (mitochondrial, proton carrier)	TGCACCACAG [G/A] AGCCATGGCG	υ Σ	A	р Н
G556u3	WIAF-11823	AF001787	1113	UCP2, uncoupling protein 2 (mitochondrial, proton carrier)	TACGGGAATC[A/G]CCGTTTTGAA	8	ъ. С	ري دي
G556u4	WIAF-11782	AF001787	386	UCP2, uncoupling protein 2 (mitochondrial, proton carrier)	ATCCTGACCA [T/C] GGTGCGGACT	E	υ	E
G561a1	WIAF-12111	HT1176	2430	2430 IDE, insulin-degrading enzyme	ACTGTGGCAT[C/A]GAGATATACT	ပ အ	Ą	н
G561u2	WIAF-12222	HT1176	3099	3099 IDE, insulin-degrading enzyme	ATATTAACTT [C/G] ATGGCTGCAA	υ Σ	_D	딘

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G562u1	WIAF-12223	HT27503	680	tumor nec 680 type 1 a	necrosis factor receptor L associated protein	CCTGTAGTGA [A/C] TCGGCCGCTG	Σ A	υ	z	H
2562112	WTAR-12224	HT27503	006	tumor necrosis	factor receptor ed protein	CGCTGCAGCG [C/A] CTGGTGGAGG	ა ე	Æ	ద	<b>64</b>
G573u1	WIAF-12199	HT28094	469		ptor 1	GGACCGCTAC [G/C] TGGCCGTGGT	ບ ຮ	υ	>	ᆈ
G573u2	WIAF-12208	HT28094	480	480 SSTR1,	somatostatín receptor 1	TGGCCGTGGT [G/A] CATCCCATCA	S	A	>	>
G573u3	WIAF-12209	HT28094	879	SSTR1,	somatostatin receptor 1	TGCAGCTGGT [T/C]AACGTGTTTG	E S	ŭ	>	>
G574u1	WIAF-11822	HT4058	1054	SSTR5,	somatostatin receptor 5	GCCACGGAGC [C/T] GCGTCCAGAC	υ Σ	E-1	Δ,	ᆈ
G575u1	WIAF-12200	HT28095	99	SSTR3,	somatostatin receptor 3	ACGTGTCGGC [G/A] GGCCCAAGCC	Ω Ω	Æ	⋖	Æ
G575u2	WIAF-12217	HT28095	453	453 SSTR3,	somatostatin receptor 3	CCACCCGCTC [G/A] GCCCGCTGGC	ω Ω	A	Ω	တ
G585u1	WIAF-12204	HT1022	1133	PYGL, p liver (H storage	PYGL, phosphorylase, glycogen; liver (Hers disease, glycogen 1133 storage disease type VI)	AGCTGAATGA [T/C] ACTCACCCTC	ν Ε-	Ü	Д	А
6585112	WIAF-12205	HT1022	1988	PYGL, liver ( storage	PYGL, phosphorylase, glycogen; liver (Hers disease, glycogen storage disease type VI)	AGCTGATCAC [T/C] TCAGTGGCAG	S	Ü	단	E-1
G585u3	WIAF-12225	HT1022	1883	PYGL, liver ( storage	PYGL, phosphorylase, glycogen; liver (Hers disease, glycogen storage disease type VI)	IGTACAACCG [C/T] ATTAAGAAAG	S S	E-1	ద	ద
G585u4	WIAF-12226	HT1022	2037	PYGL, liver ( storage	PYGL, phosphorylase, glycogen; liver (Hers disease, glycogen storage disease type VI)	aagcaagttg [a/g] aagtcatctt	M	ڻ د	×	[2]
G585u5	WIAF-12231	HT1022	1387	PYGL, 1 liver (I	PYGL, phosphorylase, glycogen; liver (Hers disease, glycogen 1387 storage disease type VI)	GAIGTGGACC [C/G] TCTGAGAAGG	υ Σ	D)	Д-	요
G586a1	WIAF-12112	HT1878	2410	2410 PFKM,	phosphofructokinase, muscle CCGGGGAAGC[T/G]GCCGTCTAAA	CCGGGGAAGC [T/G] GCCGTCTAAA	<u>⊱</u>	<u> </u>	A.	A

G586u2 WIAF-12206 G586u3 WIAF-12207 G586u4 WIAF-12227 G586u5 WIAF-12228	HT1878						
		375	PFKM, phosphofructokinase, muscle	muscle GGACGACTCC[G/A]AGCTGCCTAC	M G A	ద	Q
	7 HT1878	322	322 PFKM, phosphofructokinase, muscle TGGGAGGCAC[G/A]GTGATTGGAA	IGGGAGGCAC [G/A] GTGATTGGAA	ა ე	Ð	E
	7 HT1878	334	PFKM, phosphofructokinase, muscle	muscle TGATTGGAAG[I/C]GCCGGTGCA	S H	ß	ഗ
	HT187	408		CGTGGGATCA [C/G] CAATCTCTGT	უ ე დ	H	ß
G586n6 WIAF-12235	HT187	717	717 PFKM, phosphofructokinase, muscle	muscle CACTGTGGAT[A/G]CCTGGCCCTT	м М		Ü
		366	hofructokinase, liver	ATGGCAGCCT [T/C] ACAGGTGCCA	S	1	긔
	0 139211	1327	CPTIA, carnitine 1327 palmitoyltransferase I, liver	CAGCGTTCTT [C/T] GTGACGTTAG	S S	Ĺτι	[24
G589u2 WIAF-12215	5 1.39211	2080	CPT1A, carnitine 2080 palmitoyltransferase I, liver	AATATCTCGC [T/C] GTGGAGTCCC	ت اط	A	Æ
	6 1139211	679	CPT1A, carnitine 679 palmitoyltransferase I, liver	ACTTCAAACG [G/T] ATGACAGCAC	S S	<u>~</u>	ρĸ
		1844	CPT1A, carnitine 1844 palmitoyltransferase I, liver	CCTCACATAC [G/C] AGGCCTCCAT	υ υ Σ	田	Q
			NSMAF, neutral sphingomyelinase (N-SMase) activation associated factor	TCCGGGATCT[C/T]AGTAAGCCAG	El C)	1	д
		2000	NSMAF, neutral sphingomyelinase (N-SMase) activation associated factor	aagtatatca [t/G] tttcaaatat	E E	[I4	Λ
G352uz MIAF-11834	96 X		NSMAF, neutral sphingomyelinase (N-SMase) activation associated 1673 factor	GTAGCCATGC [T/C] TACGCAAATC	U H E	니	Д

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	•			NSWAF, neutral sphingomyelinase (N-SWase) activation associated					
G592u4	WIAF-11784	X96586	1889		CACGAGCACT [A/G] TAAAATCCAC	Σ	ტ	×	U
G592u5	WIAF-11798	X96586	1677	NSMAF, neutral sphingomyelinase (N-SMase) activation associated factor	CCATGCTTAC [G/A] CAAATCTTGG	<u>ი</u>	4	E	F
G592u6	WIAF-11799	X96586	2429	NSWAF, neutral sphingomyelinase (N-SMase) activation associated factor	TGCCATTCAG [G/C] GATTGTATGT	<u>υ</u>	υ	_O	4
G592a7	WIAF-13156	X96586	2205	NSMAF, neutral sphingomyelinase (N-SMage) activation associated factor	ATTCTGCATC [G/A] TGGGACTCTA	დ დ	ব	w	S
G594u1	WIAF-10065	HT3921	1153	1153 annexin V, alt. transcript 2	TTGTGAAATC [T/A] ATTCGAAGTA	Ω ⊟	Ø	ß	ß
G594u2	WIAF-10098	HT3921	567	annexin V, alt. transcript 2	CGAAGTAATG [C/T] TCAGCGCCAG	υ Σ	E	Æ	Þ
G594u3	WIAF-10099	HT3921	774	annexin V, alt. transcript 2	ATTGCTTCAA [G/C] GACACCTGAA	υ Σ	ပ	ω.	EH
G594a4	WIAF-10505	HT3921	424	annexin V, alt. transcript 2	GAGTAGTCGC [C/T] ATGGCACAGG	ט	EH		
G594a5	WIAF-13123	HT3921	571	annexin V, alt. transcript 2	GTAATGCTCA [G/C] CGCCAGGAAA	IJ Œ	υ	Q	н
G595u1	WIAF-12203	HT27983	1008	NRIP1, nuclear receptor 1008 interacting protein 1	TGCAAGATTA [C/T] AGGCTGTTGC	N	E	Ø	*
G595u2	WIAF-12220	HT27983	785	NRIP1, nuclear receptor interacting protein 1	CCCTCAGTCA [T/C] GATTCTTTAA	E	υ	Ħ	н
G595u3	WIAF-12232	HT27983	1231	NRIP1, nuclear receptor 1231 interacting protein 1	GTTGGCAGTT [A/T] CCAGCTCCCA	Σ	⊟	×	ഥ
G595u4	WIAF-12261	HT27983	2048	NRIP1, nuclear receptor 2048 interacting protein 1	GCAGTACTCA [G/A] TCTGAAAAGC	Ω D		Ø	Q
G595u5	WIAF-12274	HT27983	2376	NRIP1, nuclear receptor interacting protein 1	TCCTGAACCA [G/T] GGCTTTCTGG	ع ق	E	ტ	×
G595u6	WIAF-12275	HT27983	3498	NRIP1, nuclear receptor 3498 interacting protein 1	ACTATATTAC [A/G] TGCTTCAAAA	M	ტ	Σ	>
G595u7	WIAF-12276	HT27983	3671	NRIP1, nuclear receptor interacting protein 1	ACAATAGCCA [T/C] ATGGGAAATA	ST	U	田	Ħ
G595u8	WIAF-12294	HT27983	2020	NRIP1, nuclear receptor 2020 interacting protein 1	ATCAAATGGA [A/G] TTCCCCACCA	M	<u> </u>	_ z	ഗ

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2595119	WIAF-12295	HT27983	3140	NRIP1, nuclear receptor 3140 interacting protein 1	ATTTGTCCCC[G/A]CACAGAAGTA	S	A A	<u></u> 전
G596u1	WIAF-10144	HT3537	3299 PC,	ylase	TGCGGTCCAT [C/T] TTGGTCAAGG	၁ ဒ	Ft.	H H
G596u2	WIAF-10158	HT3537	2662 PC,	pyruvate carboxylase	ACCAACCTGC [A/C] CTTCCAGGCC	Ą	D.	
G596u3	WIAF-10159	HT3537	2156 PC,	pyruvate carboxylase	CCATCTCATA[C/A]ACGGGCGACG	O Z	A	*
G598a1	WIAF-12118	HT48666	55 85 58 55	HERC1, hect (homologous to the E6-AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1	GGGACCTATG [C/T] TGATAAACTG	υ Σ	- L	\delta
G598u2	WIAF-12236	HT48666	4456	HERC1, hect (homologous to the E6-AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain	CCTGTTAATA [T/C] TAGGAGTAAG	E S	υ υ	그
G598u3	WIAF-12237	HT48666	6356	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 6356 (RLD) 1	GGTAATGAAG [G/T] CACGTGTGTT	υ Σ	ы	۸ و
G598u4	WIAF-12240	HT48666	12219	HERCI, hect (homologous to the E6APP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain (RLD) 1	GTACCTTTGT [C/T]ATCCAGGCCA	<u>ი</u>	F	Λ Λ
G598u5	WIAF-12241	HT48666	12480	HERCI, hect (homologous to the E6. AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1	CCAGGCAGAT [C/G] GAGGCCTTAC	υ Σ	ย	Σ Η
G598u6	WIAF-12244	HT48666	12975	HERC1, hect (homologous to the E6-AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1	GAGTAATCAT [T/A] GAAGATGTGG	ω Ε-ι	A	H
G598u7	WIAF-12245	HT48666	1424	HERC1, hect (homologous to the E6. AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1	TCCAATAATC [A/T] GTCAACTTTA	Æ	F	Fi O

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HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain 7635 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain 9189 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain 10119 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain 11109 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain 13513 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus)
HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 9189 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 10119 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 11109 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 13513 (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus)
HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus)
HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus)
HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain (RLD) 1  HERCI, hect (homologous to the E6- AP (UBE3A) carboxyl terminus)
HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus)
domain and RCC1 (CHC1)-like domain ACCATCACAG[A/G]GATGIGCCAG

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carboxyl terminus) RCCI (CHCI)-like c	AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain	AP (UBE3A) of domain and FHT48666 1098 (RLD) 1
C U U	HERC1, hect (homologous to the E6APP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 6079 (RLD) 1	HERC1, hect AP (UBE3A) codomain and R HT48666 6079 (RLD) 1
t (homologous to the E6 carboxyl terminus) RCC1 (CHC1)-like domain	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 9551 (RLD) 1	HERC1, hec AP (UBE3A) domain and (RLD) 1
# # U	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 666 (RLD) 1	HERC1, hect AP (UBE3A) ca domain and RC HT48666 666 (RLD) 1
- उत्स	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 882 (RLD) 1	HERC1, hect AP (UBE3A) can domain and RCI HT48666 882 (RLD) 1
	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 893 (RLD) 1	HERC1, AP (UBE) domain a
	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 13276 (RLD) 1	HERC1, hect ( AD (UBE3A) car domain and RCC HT48666 13276 (RLD) 1
	HERC1, hect (homologous to the E6 AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like domain 6519 (RLD) 1	HERC1, hect () AP (UBE3A) carl domain and RCC HT48666 6519 (RLD) 1

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HERC1, hect (homologous to t AP (UBE3A) carboxyl terminus) domain and RCC1 (CHC1)-like d HT48666 1197 (RLD) 1
HERC1, hect (homolo AP (UBE3A) carboxyl domain and RCC1 (CHC HT48666 3595 (RLD) 1
HERC1, hec AP (UBE3A) domain and 3661 (RID) 1
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protein kinase, mitogen-activated, HT48690 1012 p38Beta (MAP kinase p38Beta)
protein kinase, mitogen-activated, 799 p38Beta (MAP kinase p38Beta)
protein kinase, HT48690 848 p38Beta (MAP kir
protein kinase, mitogen-activated,

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G612u1	WIAF-12344	HT1436	RAF1, 1513 viral	v-raf-1 murine leukemia oncogene homolog 1	TTTGCATGCA[A/G]AGAACATCAT M A	_O	м	БД
G614u1	WIAF-12267	HT321	603	BRAF, v-raf murine sarcoma viral oncogene homolog Bl	GACAGTCTAA [A/G] GAAAGCACTG MAA	<u> </u>	×	px.
G614u2	WIAF-12268	HT321	2282	BRAF, v-raf murine sarcoma viral oncogene homolog Bl	CCAAACAGAG [G/A] ATTTTAGTCT M G	A	Д	z
G614u3	WIAF-12299	HT321	973	BRAF, v-raf murine sarcoma viral oncogene homolog B1	AGGAAGAGGC[G/A]TCCTTAGCAG S G	<b>4</b>	Ø	A
G616u1	WIAF-12253	HT48746	498	TRAF-interacting protein (I-TRAF)	AAGAAGACAA [G/T] AGGTTTCTTC N G	<u>[+ </u>	ы	*
G616u2	WIAF-12269	HT48746	1338	1338 TRAF-interacting protein (I-TRAF)	GCATATACCT [C/G] GAGTATGTGA M C	ט	DK.	U
G616u3	WIAF-12285	HT48746	377	377 TRAF-interacting protein (I-TRAF)	ATAACAATTA [T/C] GGCTGTGTCC S T	υ H	>-	>-
G616u4	WIAF-12288	HT48746	1032	1032 TRAF-interacting protein (I-TRAF)	TGAAATTCAG [G/A] GAATTGACCC M G	۵ ک	ტ	ద
G617u1	WIAF-12256	HT1614	52	PPPICA, protein phosphatase 1, catalytic subunit, alpha isoform	GAAGCTCAAC [C/T] TGGACTCGAT S (	H ت	- Ц	ı
E	# + + + + + + + + + + + + + + + + + + +	HT1614	792	PPPICA, protein phosphatase 1,792 catalytic subunit, alpha isoform	AAGACGGCTA[C/T]GAGTTCTTTG S	E4 O	>1	Y
G61/u2	WIAK-12270	HT27508	1598	protein phosphatase, 2A B56-alpha subunit	CATTGAACCA [A/C] CACAGTTCAA	C A	H	Д
Gerour	WIRE-12271	HT27508	1135	protein phosphatase, 2A B56-alpha subunit	ATCAGAAATT [C/T] GTACAACAGC	H ن	<u>[24</u>	[z ₁
ZDO TOS				ERCC6, excision repair cross-			1	
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G62u1	WIAF-10369	HT0855	214	9		Ī		_
				ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group	M GCCARROTT (7) ITT HOMBARA	<u>U</u>	[T4	н
G62u2	WIAF-10370	HT0855	926	9				-
				ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group	AGCACGGACA [C/T] GCAGGCCCGG	ت ب	H	Σ
G62u3	WIAF-10428	HT0855	2904 6	10				

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				ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group				>	
G62u4	WIAF-10430	HT0855	3368	9	TGACCCTCAC [A/G] TGAGTAGTAA	E E	5	E	Ţ
G62115	WIAF-10451	HT0855	1376	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	TTCTGGGGAA [G/A]AAGCTGAAGC	ڻ ح	A.	M	×
G62u6	WIAF-10452	HT0855	E C C C A 2716 6	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	TAAGCATTGC [A/G] GAGACGCCAA	ع ع	0	ద	_O
G62117	WIAE-10453	HT0855	3967	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	CCCTGAAAGC [A/C] CTGAGGCTCT	ر ا	ن	Æ	Æ
811695	WTAF-10454	HT0855	4016	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	TGGTGTTCCC [A/G] CCTGGACTGG	Z Z	ŭ	<u> </u>	ď
6n299	WIAE-10455	HT0855	3979	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	TGAGGCTCTC [T/C] CGTCAGCGGT	S E	ט	S	တ
G62m10	WIRE-10456	HT0855	3729 6 Q. Q.	ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group 6	GACGCCAAGT [T/G] TGAAGGAACT	<u>F</u>	ט	ഥ	ن
G62u11	WIAE-10476	HT0855	1275 6	ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group 6	TCTGGAGATG [G/A] TACTGACTAT	<b>ʊ</b>	A.	ರ	Д
G621112	WIAE-10477	HT0855	2017	ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group	TGATCITGGA [C/T] GAAGGACACA	න ව	EH	Ω	Д
G62u13	WIAF-10479	HT0855	EI CC CC 3265 6	ERCC6, excision repair cross-complementing rodent repair deficiency, complementation group 66	CTAACATATC[T/C]GTAAATGATG	S	<u></u> υ	ω	w

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GGGCACCTGC [A/G] GGAAGCTTCT	TATCATGGAA [T/A] TAGATGACAC	CCTCATGTTA [C/G] ACGGCGCACC	ttttatgatg [a/g]atgtctgcga	TTCATGGACA[A/G]TATACAGATT	CATTCTGGAG [A/G] ATTACTAGGA	AGGGGTATGA [T/A] CACAAAGCAA	CCAATTAITG[C/T]GGAGAGITTG	GATCTTATAT [G/T] TAGAGCCCAT	AAAGATGCAG [A/G] TCTGAACTCT	CGATGGGAAC [G/T] CCCCATCCTT	CGCCCCATCC[T/c]TTGGTTTACT	ACTTAAAGGA [T/C]ATTGCAGGAG	TAAAGATGTG [C/T] TTGGACATCT
ERCC6, excision repair cross- complementing rodent repair deficiency, complementation group 6	PPP2CB, protein phosphatase 2 (formerly 2A), catalytic subunit, 1256 beta isoform	PPP2CB, protein phosphatase 2 (formerly 2A), catalytic subunit, beta isoform	PPP2CB, protein phosphatase 2 (formerly 2A), catalytic subunit, 819 beta isoform	PPPICB, protein phosphatase 1, 459 catalytic subunit, beta isoform	PPP2R2A, protein phosphatase 2 (formerly 2A), regulatory subunit B (PR 52), alpha isoform	PPPICC, protein phosphatase 1, 1104 catalytic subunit, gamma isoform	PPPICC, protein phosphatase 1, 973 catalytic subunit, gamma isoform	PPPICC, protein phosphatase 1, 888 catalytic subunit, gamma isoform	protein phosphatase 2A, 130 kDa regulatory subunit	protein phosphatase 2A, 130 kDa regulatory subunit	protein phosphatase 2A, 130 kDa regulatory subunit	protein phore	protein phosphatase 2A, 130 kDa 1200 regulatory subunit
H 4317	1256	PPP2C   (form   1326 beta	819	459	227	1104	973	88 8	704	1015	1024	837	1200
HT0855	HT1943	HT1943	HT1943	HT3979	HT1961	HT2780	HT2780	HT2780	HT5086	HT5086	HT5086	HT5086	HT5086
WIAF-10481	WIAF-12116	WIAF-12117	WIAF-12239	WIAF-12260	WIAF-12266	WIAF-12104	WIAF-12105	WTAF-12311	WTAR-12103	WIAF-12106	WT&R-12107	WTAR-12108	WIAF-12325
G62u14		G620a2	G620u3	G623u1	G625u1	G628al	G628a2	5.18.09	G02000	G602092		10000000000000000000000000000000000000	G630u5

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9110898	WIAF-12326	HT5086	2810 r	protein phosphatase 2A, 130 KDa regulatory subunit	ATGTTCAGGG [C/T] TGCAGGGGGA	υ Σ	H	⋖	Þ
3630u7	WIAF-12351	HT5086	512 r	protein phosphatase 2A, 130 kDa regulatory subunit	ATTATGGCAG [C/T] AACTTACAGA	υ Σ	_ E	Æ	>
0110000	WTAR-12352	HT5086	703 r	protein phosphatase 2A, 130 kDa requlatory subunit	CAAAGATGCA [G/A] ATCTGAACTC	<u>დ</u>	Æ	Д	z
0000000	CHOCK ERFE	780 HH		osphatase 2A, 130 kDa subunit	ACCTTTGTCT [C/T] ATAGAAACTC	υ Σ	단	田	≽ı
G63009	WLAF -12333	000		ke growth factor	TGCAAGTGGC [C/T] AACACCACCA		<u></u>	Ø	Ą
G634u1	WIAF-11825	X04434	T C077	remp insulin-like growth factor					
G634u2	WIAF-11826	X04434	2279 1	ptor	GTCATGCAAG [T/C] GGCCAACACC	E	υ	>	Æ
01.7000	WT&R-11781	X04434	1731	IGFIR, insulin-like growth factor 1 receptor	ACAAGGACGT [G/A] GAGCCCGGCA	ω υ	A	>	Λ
רח די רי סיי				sulin-like growth factor	TCCACGACGG [C/A] GAGTGCATGC	ა ე	Æ	r	υ
G634a4	WIAF-13106	X04434	848	poor			_	_	
G634a5	WIAF-13107	X04434	1089		CTTCTGCTCA [G/C] ATGCTCCAAG	υ Σ	ט	O.	H
26.24.26	WTAF-13108	X04434	2539	IGFIR, insulin-like growth factor 1 receptor	agaaggagca [g/a] atgacattcc	υ Σ	4	Δ.	z
			C	IGFIR, insulin-like growth factor	AAGIGGCCGG [A/C]ACCIGAGAAT	M A	υ	团	Ą
G634a7	WIAF-13109	A04434	000	IGF1R, insulin-like growth factor				!	
G634a8	WIAF-13111	X04434	1543	ptor	CTCCACCACC [A/T] CGTCGAAGAA	Σ	₽		ω
Q 4 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	WTAF-13112	X04434	1549	IGFIR, insulin-like growth factor 1 receptor	CACCACGICG [A/G] AGAATCGCAT	Æ	Ö	×	[34]
22.00	WTAR-13113	X04434	1596	IGFIR, insulin-like growth factor 1 receptor	CCCCTGACTA [C/T] AGGGATCTCA	S	E	>-	Þ
G645u1	WIAF-12332	HT5191	1127	retinoic acid-binding protein II	TCTGCAGACT[C/T]TTCAGGAGAG	Σ Σ	<u>[-1</u>	ㅂ	Ľt4
G645u2	WIAF-12333	HT5191	1048	1048 retinoic acid-binding protein II	AAGCATTAGA [G/A] GCCTTACAGA	S	A	[22]	回
			(	EMR1, eg mucin-lik	TAPATA (T./C.) GTGGACTAPA	E	Ü	Σ	H
G646u1	WIAF-12303	X81479	1204	r action to					_
	10 5 C L - W G T W	X81479	1919	EMR1, egf-like module containing, mucin-like, hormone receptor-like sequence 1	TTCTGCTGTG [T/G] CGCTCCATCC	E	<u>_</u>	Ü	8

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WIAF-12316	· - · · · · · · · · · · · · · · · · · ·		EMR1, egf-like module containing,				
	X81479	3 065	mucin-like, hormone receptor-like seguence 1	CTTGCCCAGA[G/T]CATGCAACTT M	ප	- F	<u>А</u>
WIAF-12317	X81479	799 s	EMR1, egf-like module containing, mucin-like, hormone receptor-like sequence 1	GCACCAAGCA [G/A] TGGACAGTTG M	<u></u> 0	K	S
WIAF-12318	X81479	558	EMR1, egf-like module containing, mucin-like, hormone receptor-like sequence 1	TGAAGACGTG [A/G] ATGAATGTGC	A	U	D Z
WIAF-12334	X81479	207	EMR1, egf-like module containing, mucin-like, hormone receptor-like 207 sequence 1	TTACTATTGC [A/G] CTTGCAAACA	ĸţ	Ü	T A
WIAF-12335	X81479	458	EMR1, mucin-1 sequenc	TCACCAGCAG [G/C] GTCTGCCCTG M	<b></b>	Ü	<u> </u>
WIAF-12336	X81479	1308	EMR1, mucin-1 sequenc	CTCAGCAAAT [G/A] TCACTCCGGC M	ro O	A	H
WIAF-12337	X81479	1285	EMR1, mucin-1 sequenc	ACACTGGCAT [C/T] TTTTGGAAA M	υ _	E-I	Ω [± ^t
2338	X81479	2026	EMR1, mucin-1 sequenc	GACAACAAGA[C/T]GGGCTGCGCC M	U U	E	Σ H
2339	HT5190	174	RARA, retinoic acid receptor, alpha	TGCCTCCCTA [C/T] GCCTTCTTCT	U FO	F	A A
WIAF-13332	HT0070	469	retinoic acid receptor, beta	AACGTGAGCC [A/G] GGAGCAGCGT	- ₹	ŋ	1
WIAF-13333	HT0070	532	retinoic acid receptor, beta	ATTGTTTTA [A/G] GGTGAGAAAT	Æ	ರ	1
WIAF-12323	X52773	862	RXRA, retinoid X receptor,	CTCGCCGAAC [G/A] ACCCTGTCAC	<u>ი</u>	Ø	Ω Ω
WIAF-12341	X52773	102	retinoid X receptor,	TCCTGCCGCT [C/T] GATTTCTCCA	D	ĘH	1
	MIAF-12335 MIAF-12336 WIAF-12338 WIAF-12339 WIAF-13332 WIAF-13333 WIAF-13333	X8147 X8147 X8147 X8147 X8146 HT515 HT516 HT00' HT00' X527'	X81479 1: X81479 1: X81479 2 HT5190 HT0070 HT0070  HT0070	EWR1, egf-like module containulus	EWR1, egf-like module containing,   mucin-like, hormone receptor-like   TCACCAGCAG[G/C]GTCTGCCCTG   Mucin-like, hormone receptor-like   TCACCAGCAG[G/C]GTCTGCCCTG   Mucin-like, hormone receptor-like   CTCAGCAAAT[G/A]TCACTCCGGC   Mucin-like, hormone receptor-like   ACACTGGCAT[C/T]TTTTGGAAA   Mucin-like, hormone receptor-like   ACACTGGCAT[C/T]TTTTTGGAAA   Mucin-like, hormone receptor-like   ACACTGGCAT[C/T]TTTTTGGAAA   Mucin-like, hormone receptor-like   ACACTGGCAT[C/T]TTTTTGGAAA   Mucin-like, hormone receptor-like   ACACTGGCAT[C/T]GCCTTCTCT   Albha   ATTGTTTTTA[A]GGCTGCGCC   ACCTGCTGTGCCC   ACCTGCTGTCCCA   ACCTGCTGTCCCA   ACCTGCTGTGCCCCA   ACCTGCTGTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCTCCCA   ACCTGCTGTCCCA   ACCTGCTGTCCCA   ACCTGCTGTCCCA   ACCTGCTGTCCA   ACCTGCTGTCA   ACCTGCTGTCA   ACCTGCTGTCA   ACCTGCTGTCA   ACCTGTCTCA   ACCTGCTGTCA   ACCTGCTGTCA   ACCTGCTGTCA   ACCTGTCTCA   ACCTGCTGTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCTCA   ACCTGTCA   ACC	EMR1, egf-like module containing, mucin-like, hormone receptor-like module containing, mucin-like, hormone receptor-like mucin-like, hormone receptor-like carcacacacacacacacacacacacacacacacacaca	EWR1, egf-like module containing, mucin-like, hormone receptor-like   TCACCAGCAG[G/C]GTCTGCCCTG   M G   C

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G650u4 WIAF-12349 X52773 G653al WIAF-13326 HT1458 G655al WIAF-13327 J05252 G655a2 WIAF-13334 J05252 G658u2 WIAF-13335 J02943 G658u3 WIAF-13407 J02943 G658u4 WIAF-13409 J02943 G668u4 WIAF-13409 J02943 G668u1 WIAF-13400 HT3157 G668u2 WIAF-13350 U53506 G668a2 WIAF-13351 U53506 G668a2 WIAF-13352 U53506 G673a1 WIAF-13352 U53506		tinoid X receptor, alpha orinoic acid receptor,  noprotein convertase  noprotein convertase  noprotein convertase  noprotein convertase  noprotein convertase  noprotein convertase  noprotein convertase  rticosteroid binding	GACAAACAGC [T/C] TTTCACCCTG  AGGAGAAAGC [T/C] CTCAAAGCAT  CCTTCAGCAA [C/T] GGGAGGAAAA  CCTATCCTTA [C/A] CCTCGGTACA  TTTCTGCTGC [C/T] GCCAACACA  TCTATGACCT [T/C] GGAGATGTGC			L & * & D
WIAF-13326 WIAF-13327 WIAF-13334 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13400		ertase ertase ending nding		H 0 0 0 H		F. A * Z A
WIAF-13327 WIAF-13334 WIAF-13335 WIAF-13409 WIAF-13409 WIAF-13409 WIAF-13400 WIAF-13400 WIAF-13400 WIAF-13400 WIAF-13400 WIAF-13350 WIAF-13350 WIAF-13350	П .	ertase ertase oding		U U H		Z * Z
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WIAF-11856 WIAF-13407 WIAF-13409 WIAF-13400 WIAF-13401 WIAF-13350 WIAF-13351 WIAF-13336	176	rticosteroid binding rticosteroid binding	CTATGACCT [T/C] GGAGATGTGC	E-I		ㅁ
WIAF-13407 WIAF-13408 WIAF-13409 WIAF-13401 WIAF-13350 WIAF-13351 WIAF-13336	1771	rticosteroid binding	777 E7 5 47 47 E7 E1 5 4 5 E 47 E 47 E 47 E 47 E 47 E 47 E 4	_		
WIAF-13408 WIAF-13409 WIAF-13400 WIAF-13401 WIAF-13350 WIAF-13351 WIAF-13358		globulin	CTTCATGAC [T/6] CAGAGCICCC	T E		A
WIAF-13409 WIAF-13400 WIAF-13401 WIAF-13350 WIAF-13351 WIAF-13328 WIAF-13336	773	CBG, corticosteroid binding globulin	TTCATGACTC [A/G] GAGCTCCCCT	S A S		တ
WIAF-1336 WIAF-13350 WIAF-13351 WIAF-13358 WIAF-13336		CBG, corticosteroid binding	TCACCCAGGA [C/T] GCCCAGCTGA	SCT	D	Д
MIAF-13401 HT31 WIAF-13350 U535 WIAF-13352 U535 WIAF-13328 M574	-	yroid peroxidase	CGCCACGCGC [G/A] CCTGCGGCCT	Ö	4	Æ
WIAF-13350 U535 WIAF-13351 U535 WIAF-13352 U535 WIAF-13328 M574	7 1282 TPO,	thyroid peroxidase	GGCCGCCCA [G/C] CGAGGTCCCC	D M	ω	H
WIAF-13351 U535 WIAF-13328 W574 WIAF-13336 M574	350	DIO2, deiodinase, iodothyronine,	TCGATGCCTA [C/A] AAACAGGTGA	N O	<i>&gt;</i> √	*
WIAF-13352 U535 WIAF-13328 M574	354	DIO2, deiodinase, iodothyronine, 54 type II	TGCCTACAAA [C/A] AGGTGAAATT	M C A	O)	- *
WIAF-13328 M574		DIO2, deiodinase, iodothyronine,	TGTCTCCAGT [A/G] CAGAAGGAGG	M A	EH U	₹.
WIRF-13336 M574		Human ret proto-oncogene mRNA for 1723 tyrosine kinase.	CGAGCCTGGG [G/A] AGCCCCGGGG	Σ Q	A	×
		Human ret proto-oncogene mRNA for 1186 tyrosine kinase.	GGCTCGCCGA [T/A] TTGCCCAGAT	E E	A F	
WIAF-13337		Human ret proto-oncogene mRNA for 1227 tyrosine kinase.	ACTGCCAGGC [G/A] TTCAGTGGCA	S G	A	Æ
G673a4 WIAF-13338 M57464		Human ret proto-oncogene mRNA for 2118 tyrosine kinase.	TTGGAAAAC[T/A]CTAGGAGAAG	S F	A T	
G673a5 WIAF-13339 M57464		Human ret proto-oncogene mRNA for 2238 tyrosine kinase.	CGAGTGAGCT [T/G] CGAGACCTGC	E S	E G	므

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G678al	WIAF-13353	D49492	1439 f	GDF10, growth differentiation	TCGGCTGGAA [T/A] GAATGGATAA	T.	A	z	×
G68u1	WIAF-10434	HT1115	1214 6	ERCC3, excision repair cross- complementing rodent repair deficiency, complementation group 3 (xeroderma pigmentosum group B	CTGTGGAGCA [G/A] TGGAAAGCCC	න ග	4	Q	Ø
G68u2	WIAF-10435	HT1115	1155	ERCC3, excision repair cross- complementing rodent repair deficiency, complementation group 3 (xeroderma pigmentosum group B 1155 complementing)	TGTGACTGCT [G/C] CATGCACTGT	υ Σ	Ü	4	Сч
G68u3	WIAF-10436	HT1115	1327	ERCC3, excision repair cross- complementing rodent repair deficiency, complementation group 3 (xeroderma pigmentosum group B 1327 complementing)	AGCACCTACT [C/T] CATGCTGGGC	υ Σ	E E	တ	<u>Γ</u> τι
G68u4	WIAF-10461	HT1115	926	ERCC3, excision repair cross- complementing rodent repair deficiency, complementation group 3 (xeroderma pigmentosum group B 926 complementing)	AGGAAATGAT [T/C] GAGGAACTCC	© Ε-	<u>ن</u>	н	Н
G68u5	WIAE-10464	HT1115	1430	ERCC3, excision repair cross- complementing rodent repair deficiency, complementation group 3 (xeroderma pigmentosum group B	AAGTGCACAC [C/T] ATACCAGCCA	υ υ	E+	E	H
G684a1	WIAF-13359	X51801	712	BMP7, bone morphogenetic protein 712 7 (osteogenic protein 1)	GTTTATCAGG [T/G] GCTCCAGGAG	E E	r r	>	_O
G684a2	WIAF-13360	X51801	719	BMP7, bone morphogenetic protein 719 7 (osteogenic protein 1)	AGGTGCTCCA [G/A] GAGCACTTGG	ω υ	A	Q	ø
G684a3	WIAF-13361	X51801	BM.	BMP7, bone morphogenetic protein 7 (osteogenic protein 1)	GGCTGGCTGG [T/G] GTTTGACATC	E	<u>.</u>	⊳	Ð

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MMP7, bone morphogenetic protein   X51801   658 7 (osteogenic protein 1)   X51801   1421 7 (osteogenic protein 1)   General September   1421 7   General September   1421 7   General September   1421 7   General September   1421 7   General September   1422 7   General September   1422 8   General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1688 General September   1767 Centium-sensing receptor    G684a4	WIAF-13362	X51801	862	BMP7, bone morphogenetic protein 862 7 (osteogenic protein 1)	GGCCTGCAGC [T/G] CTCGGTGGAG	T	D)	FI	EK.	
MIAF-13834   X51801   1421 7 (osteogenic protein 1)   0   0   0   0   0   0   0   0   0	G684a5	WIAF-13363	X51801	658	c protein	ATCTACAAGG[A/G]CTACATCCGG	Æ	Ŋ	Д	Ü
MIAF-13329   D89675   S82   Protein receptor, type IB	G684u6	WIAF-13834	X51801		bone morphogenetic protein teogenic protein 1)	GCCACTAGCT [C/T] CTCCGAGAAT	<u>ن</u> -	E	1	
MIAF-13330   D89675   920 protein receptor, type IB	G685a1	WIAF-13329		882	bone morphogenetic receptor, type IB	GTTCCCTTTA [T/G]GATTATCTGA	F	_O	X	*
MIAF-13331   D89675   770 protein receptor, type IB	G685a2	WIAF-13330	D89675	920	H	GCTAAATCAA [T/C]GCTGAAGTTA	E E	υ	Σ	₽
MIAF-13340   D89675   1303 protein receptor, type IB	G685a3	WIAF-13331	D89675	770	H H	TATCAGACAG [T/G] GTTGATGAGG	E E	Ö	>	U
WIAF-13341   D89675   1372 protein receptor, type IB	G685a4	WIAF-13340	D89675	1303	Н	TCCTTATCAT [G/A] ACCTAGTGCC	Σ Ω	4		z
MIAF-13342   D89675   1173 protein receptor, type IB	G685a5	WIAF-13341	D89675		H	GTTACGCCCC[T/G]CAFTCCCAAA	Σ	U	w	Æ
WIAF-13816   Z48923   2705 (serine/threonine kinase)   AIAF-13816   Z48923   2705 (serine/threonine kinase)   AIAF-13817   Z48923   Z749 (serine/threonine kinase)   AIAF-13343   HT1455   G26 (CALB1, Calbindin 1, (28kD)   AIAF-11839   HT27700   1075 (calcium-sensing receptor   WIAF-11840   HT27700   1551 (calcium-sensing receptor   WIAF-11841   HT27700   1688 (calcium-sensing receptor   WIAF-11842   HT27700   1698 (calcium-sensing receptor   MIAF-11858   HT27700   1698 (calcium-sensing receptor   MIAF-11858   HT27700   1698 (calcium-sensing receptor   MIAF-11858   HT27700   1698 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (calcium-sensing receptor   MIAF-11859   HT27700   1689 (ca	G685a6	WIAF-13342			н	TGTTGGACGA[G/A]AGCTTGAACA	<u>დ</u>	4	ш	ы
WIAF-13817         Z48923         2749         Erceptor, type II           WIAF-13843         HT1455         626         CALBI, calbindin 1, (28kD)           WIAF-11839         HT27700         1075         calcium-sensing receptor           WIAF-11840         HT27700         1551         calcium-sensing receptor           WIAF-11841         HT27700         1688         calcium-sensing receptor           WIAF-11842         HT27700         1698         calcium-sensing receptor           WIAF-11858         HT27700         1698         calcium-sensing receptor           WIAF-11858         HT27700         1767         calcium-sensing receptor           WIAF-11859         HT27700         1689         calcium-sensing receptor	G686u1	WIAF-13816	Z48923	2705	BMPR2, bone morphogenetic receptor, type II (serine/threonine kinase)	aaatttggca [g/a] caagcacaaa	D E	4	ത	z
WIAP-13343         HT1455         626 CALB1, calbindin 1, (28kD)           WIAF-11839         HT27700         1075 calcium-sensing receptor           WIAF-11840         HT27700         1551 calcium-sensing receptor           WIAF-11841         HT27700         1688 calcium-sensing receptor           WIAF-11842         HT27700         1698 calcium-sensing receptor           WIAF-11858         HT27700         1767 calcium-sensing receptor           WIAF-11859         HT27700         1689 calcium-sensing receptor	G686u2	WIAF-13817	Z48923	2749	BMPR2, bone morphogenetic protein receptor, type II (serine/threonine kinase)	TGGAGTTGCC [A/T] AGATGAATAC	z z	E	M	-*
WIAF-11839         HT27700         1075 calcium-sensing receptor           WIAF-11840         HT27700         1551 calcium-sensing receptor           WIAF-11841         HT27700         1688 calcium-sensing receptor           WIAF-11842         HT27700         1698 calcium-sensing receptor           WIAF-11858         HT27700         1767 calcium-sensing receptor           WIAF-11859         HT27700         1689 calcium-sensing receptor	G687al	WIAF-13343	HT1455	626	calbindin 1,	ATGATCAGGA[C/T]GGCAATGGAT	S C	F	Ω	Д
WIAF-11840         HT27700         1551 calcium-sensing receptor           WIAF-11841         HT27700         1698 calcium-sensing receptor           WIAF-11842         HT27700         1698 calcium-sensing receptor           WIAF-11858         HT27700         1767 calcium-sensing receptor           WIAF-11859         HT27700         1689 calcium-sensing receptor	G696u1	WIAF-11839	HT27700	1075	calcium-sensing receptor	GGGCACAATT [G/C] CAGCTGATGA		ບ	A	Д
WIAF-11841         HT27700         1688 calcium-sensing receptor           WIAF-11842         HT27700         1698 calcium-sensing receptor           WIAF-11858         HT27700         1767 calcium-sensing receptor           WIAF-11859         HT27700         1689 calcium-sensing receptor	G696u2	WIAF-11840	HT27700	1551		TACCTGTGGA [C/T] ACCTTTCTGA		E-1 1		
WIAF-11858         HT27700         1698 calcium-sensing receptor           WIAF-11859         HT27700         1767 calcium-sensing receptor           WIAF-11859         HT27700         1689 calcium-sensing receptor	G696u3	WIAF-11841	HT27700	1688		TTACGGATAT [C/T] CTACAATGTG			3 3	7 F
WIAF-11859 HT27700 1689 calcium-sensing receptor	G696u4	WIAF-11842	HT27700	1698	calcium-sensing receptor	CCTACAATGT [G/T] TACTTAGCAG	-	EH E	> ⊦	>  -
WIAF-11859 H12//UU 1009 CGICLUM-SCHSING ICCCECT	G696u5	WIAF-11858	HT27700	1767	calcium-sensing receptor	GGAGAGGCT [C/T] TTCACCAATG TACGGATATC [C/T] TACAATGTGT	v s		3 C	ı s
2541 calcium-sensing receptor	G696u7	WIAF-11860	HT27700	2541	calcium-sensing receptor	TCGIGCICIG [C/T] ATCICAIGCA			ပ	υ

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G696u8	WIAF-11861	HT27700	2581 c		TGTCCTCCTG [G/A] TGTTTGAGGC	-	€ ₹	> ;	£ ;
G69695	WIAF-11863	HT27700	3159 c	3159 calcium-sensing receptor	TCTCCCGCAA [G/C] CGGTCCAGCA	T	U	×	z
G696u10	WIAF-11872	HT27700	562 c	562 calcium-sensing receptor	TCCTATTCAT [T/A] TTGGAGTAGC	E	Æ	Ē4	н
G696u11	WIAF-11878	HT27700	2941	2941 calcium-sensing receptor	CATTCCAGCC [T/G] ATGCCAGCAC	Ή Σ	ŋ	×	Ω
G696u12	WIAF-13386	HT27700	1145	1145 calcium-sensing receptor	AGGGATATCT [G/A] CATCGACTTC	υ Σ	Æ	ပ	×
G696u13	WIAF-13395	HT27700	670	670 calcium-sensing receptor	GATATTTGCC [A/G] TAGAGGAGAT	MA	ტ	н	>
G696u14	WIAF-13396	HT27700	2243	2243 calcium-sensing receptor	TTCTGGTCCA [A/G] TGAGAACCAC	M	ъ	z	ß
G696u15	WIAF-13397	HT27700	2742	2742 calcium-sensing receptor	AGCTGGAGGA [T/C] GAGATCATCT	S F	U	А	Д
G698u1	WIAF-13547	X61598	393 (	CBP1, collagen-binding protein 1	TCAGCAACTC [G/C] ACGGCGCGCA	හ ආ	ပ	S	w
G698u2	WIAF-13549	X61598	628 CBP1,	CBP1, collagen-binding protein 1	CGGCGCCCTG [C/T] TAGTCAACGC	ა ა	F		니
G698u3	WIAF-13550	X61598	1230 CBP1,	<pre>'BP1, collagen-binding protein 1</pre>	GCGGCTCCCT [G/A] CTATTCATTG	ა დ	Æ	ᄓ	L L
G701u1	WIAF-12382	HT27657	206	CGRP type I receptor	AACGATGTTG [C/A] AGCAGGAACT	M C	Ø	Æ	되
G701u2	WIAF-12391	HT27657	841 (	CGRP type I receptor	TGGACAAATT [A/T] TACCCAGTGT	Σ	F	×	ĺΣ ₄
				COL10A1, collagen, type X, alpha 1 (Schmid metaphyseal					ľ
G704ul	WIAF-14046	X60382	1396 (	chondrodysplasia)	AGGCATTCCA [G/A] GATTCCCTGG	ড হ	4	و.	Y,
G704u2	WIAF-14070	X60382	1648	COL10A1, collagen, type X, alpha 1 (Schmid metaphyseal chondrodysplasia)	TGCCAACCAG [G/C] GGGTAACAGG	<u>ن</u> ح	ಲ	O O	pt.
C C	F. C. C. C. C. C. C. C. C. C. C. C. C. C.	00000	c c	COLLOAL, collagen, type X, alpha 1 (Schmid metaphyseal	CATACCACGT [G/C] CATGTGAAAG	ა ე	<u>.</u> <u>.</u>	>	٥
G704u3	WLAF - 140 / 1	A00.004	#70T	CHOILE Out sprasses				-	
G704u4	WIAF-14072	X60382	1582	COL10A1, collagen, type X, alpha 1 (Schmid metaphyseal chondrodysplasia)	AGTCATGCCT [G/C] AGGGTTTTAT	ڻ ح	υ	м	Ø
G705al	WIAF-13228	304177	989	COL11A1, collagen, type XI, alpha 1	AGAAGAAAC[T/A]GTGACAATGA	S	ď.	EH	H
G705a2	WIAF-13229	J04177	CC 698 1	COL11A1, collagen, type XI, alpha	TGACAATGAT [T/A] GTTGATTGTA	S	A	н	н
G705a3	WIAF-13230	J04177	888	COL11A1, collagen, type XI, alpha 1	TAGTCCAGAC [T/A] GTGACTCTTC	E	Æ	υ	တ
G705a4	WIAF-13231	004177	894 1	COL11A1, collagen, type XI, alpha 1	AGACTGTGAC [T/A] CTTCAGCACC	E X	Æ	ß	E
G705a5	WIAF-13232	304177	651	COL11A1, collagen, type XI, alpha	TGACGGGAAG [T/A] GGCATCGGGT	Σ	_ 4	2	ద

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620E26	WTAF-13233	77 T 2 OT.	661 1	COL11A1,	collagen,	type XI,	alpha T	TGGCATCGGG [T/A] AGCAATCAGC	E	A.	>	斑
G705a7	WIAF-13234	704177	1597 1	COL11A1,	collagen,	type XI,	alpha	CGTCCTGGCT [T/C] ACCAGGGGCT	E	Ū E.	н	Ø
G705a8	WIAF-13235	304177	2745 1	COL11A1,	collagen,	type XI,	alpha	TGGGTTTCCA [G/A] GTGCCAATGG	Σ	<u>ب</u> ق	_O	တ
G705a9	WIAF-13236	304177	4385 1	COL11A1,	collagen,	type XI,	alpha	GTCCAGAAGG [T/A] CTTCGGGGCA	S	4	Ö	೮
G705a10	WIAF-13237	J04177	4576 1	COL11A1, 1	collagen,	type XI,	alpha	GAAAAAGGTG [A/T] CCGAGGGCTC	×	F F	Д	Λ
G705a11	WIAF-13238	J04177	4306 1	COL11A1,	collagen,	type XI,	alpha	GCTAAGGGGG [A/C]AGCAGGTGCA	Σ	ن لا	(H)	<₹
G705a12	WIAF-13239	304177	4837 1	COL11A1,	collagen,	type XI,	alpha	agacatactg [A/G] aggcatgcaa	Σ	A G	臼	ש
G705a13	WIAF-13240	304177	4931 1	COL11A1, 1	collagen,	type XI,	alpha 1	AACAAGACAT [C/T]GAGCATATGA	ß	EH U	н	н
G705a14	WIAF-13346	J04177	299 1	COL11A1,	collagen,	type XI,	alpha 7	AAGCACTAGA [T/G]TTTCACAATT	Σ	t l		闰
G705a15	WIAF-13347	304177	2225 1	COL11A1, 1	collagen,	type XI,	alpha	GGGAGCCTGG [G/C] CCTCCAGGTC	ß	ن ن	ט	Ŋ
G705u16	WIAF-13679	J04177	5493 1	COL11A1, 1	collagen,	type XI,	alpha	AATTGATCAA [G/A] TACCTATTGT	×	₽ 4	Þ	н
G705u17	WIAF-13700	304177	3484 1	COL11A1,	collagen,	type XI,	alpha	GGAGTTCAAG [G/A] TCCTGTTGGT	Σ	<b>₹</b>	<u></u> 0	Д
G705u18	WIAF-13709	304177		COL11A1,	collagen,	type XI,	alpha	GAGATGTCCT [A/T] TGACAATAAT	×	4	¥	[I4
G707u1	WIAF-12363	U32169	4996	COL11A2, 2	collagen,	type XI,	alpha	TCCCCTGAGA [C/T] TCCGTGGGGC	Σ	[ <del>.</del>   U	Н	<u>Fr</u>
G707u2	WIAF-12374	U32169	3580	COL11A2, 2	collagen,	type XI,	alpha	CAATGGCGCT [G/A] ATGGCCCACA	E	<b>∢</b>	Д	z
G707u3	WIAF-12385	U32169	2059	COL11A2, 2	collagen,	type XI,	alpha	GCCTGGCTCA [G/A] ACGGACCCCC	Σ	ر ق	Ω	z
G708a1	WIAF-13354	U73778	1885	COL12A1, alpha 1	collagen,	type XII	,	GCCTCTCCTC [C/T] TGCAGAGACC	Σ	EH D	Д	귀
G708a2	WIAF-13355	U73778	3630	COL12A1,	collagen,	type XII,	,	tgttggacaa [g/a] aaatgacaac	Σ	<b>₫</b>	田	×
G708a3	WIAF-13356	U73778	3905	COL12A1, 3905 alpha 1	collagen,	type XII	,	GCTTGTTGCA [A/T] GCTGTGGCAA	Σ	A	OX .	_=
G708a4	WIAF-13357	U73778	7051	COL12A1, 7051 alpha 1	collagen,	type XII,		ATTCCACCAG [C/A] CCGGGATGTA	Σ	Z D	A	Д
G708a5	WIAF-13358	U73778	8036	COL12A1, 8036 alpha 1	collagen,	type XII		aagaagtaaa [g/a] acattatttt	ω	ڻ ل	<u>×</u>	ᄶ

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G708a6	WIAF-13364	U73778	1461	COLIZAI, 1461 alpha 1	collagen,	type All,	TGGCTCCTAT [A/T] GCATTGGGAT	Æ	H	S	U
G708a7	WIAF-13365	U73778	2344	COL12A1, 2344 alpha 1	collagen,	type XII,	ATTACTTGGA [C/T] TCAAGCTCCA	ರ ಶ	E	H	н
G708a8	WIAF-13366	U73778	5207	COL12A1, 5207 alpha 1	collagen,	type XII,	CAGATAAGAT [G/A] GAGACCATCT	۳ ع	A	Σ	н
G708a9	WIAF-13367	073778	6592	COL12A1, alpha 1	collagen,	type XII,	GAGCCCATGG [A/T] AGCCTTTGTT	Æ	H	团	Þ
G708a10	WIAF-13368	U73778	7434	COL12A1, 7434 alpha 1	collagen,	type XII,	CCAGGATGAG [G/A] TCAAGAAGGC	უ ნ	Ą	٥	н
G708a11	WIAF-13369	U73778	9108	COL12A1, 9108 alpha 1	collagen,	type XII,	ACCTCGGGGG [C/G] TGCCTGGGCC	U E	ტ		<b>&gt;</b>
G708a12	WIAF-13370	U73778	9111	COL12A1, 9111 alpha 1	collagen,	type XII,	TCGGGGGCTG [C/T] CTGGGCCCCC	U E	E	ы	ß
G708a13	WIAF-13371	U73778	9196	COL12A1, alpha 1	collagen,	type XII,	CCCCTGGCC [G/A] TCCTGGAAAC	Σ Σ	< 4	ద	н
G708u14	WIAF-13972	U73778	3044	COL12A1, 3044 alpha 1	collagen,	type XII,	CAGTATTTGC [C/A] ACTTACAGCA	ა ა	Æ	Æ	Ą
G708u15	WIAF-13977	U73778	5853	COL12A1, alpha 1	collagen,	type XII,	TGTGACTGTA [G/C] TTCCCGTTTA	ت ت	U	>	ı,
G710u1	WIAF-12371	D38163	3082	COL19A1, alpha 1	collagen,	type XIX,	AGGAAACAAG [G/T] GCTCCATGGG	<u>ت</u>	<u>[-</u> -	ט	บ
G710112	WIAF-12388	D38163	2089	COL19A1, 2089 alpha 1	collagen,	type XIX,	TCCAGGGACT [C/T] CAGGGAATGA	υ Σ	F	<u>д</u>	S
G711u1	WIAF-12360	125286	1449	COL15A1,	collagen,	type XV, alpha	TGTGGGTCCA[A/G]GCAGTGAAGA	M A	ß	ω	ტ
G711u2	WIAF-12372	L25286	4001	COL15A1,	collagen,	type XV, alpha	ATATTCCAAT [A/G] TACTCCTTTG	Æ	_ უ	н	Σ
G711u3	WIAF-12373	L25286	3867 1	COL15A1,	collagen,	type XV, alpha	CCATTIGCAA [G/T] ATCIGICCAC	<u>ت</u>	E	Д	×
G711a4	WIAF-13372	125286	395	COL15A1,	collagen,	type XV, alpha	CCAGCAGCAC [C/T] CGTGGTGGCG	S C	E	EH	E
G711a5	WIAF-13373	125286	3101	COL15A1,	collagen,	type XV, alpha	AAGGCGACCA [G/A] GGAGCCCAGG	ა ნ	4	O)	a
G712u1	WIAF-13619	M92642	3608	COL16A1, 3608 alpha 1	collagen,	type XVI,	GGCGACCAGG [G/A] ATTTCAAGGC	Z B	A	ro l	БĦ
G712u2	WIAF-13620	M92642	4944	COL16A1, alpha 1	collagen,	type XVI,	CCATGAAAAC [C/T] ATGAAGGGGC	ა ა	EH	H	E
G712u3	WIAF-13621	M92642	4707	COL16A1, alpha 1	collagen,	type XVI,	CCAAAGGTGA [A/C] AAAGGGGACA	Æ	U	×	Д
G712u4	WIAF-13654	M92642	421	COL16A1, 421 alpha 1	collagen,	type XVI,	GCCCACGCGA [C/A] GAGTATTCCC	S S	A	p4	ద

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671713	WIRE TOOOL	250201			collagen, t	type XVI,	CHECK STRUCK (S) II AND A SECTION OF		ر	2	E
G712u6	WIAF-13656	M92642	338 a	338 alpha 1			CTCATGAAGA [A/C] GICIGCCAIC	E E		4	,
712117	WIAF-13862	M92642	3227 a.	COL16A1, CO. 3227 alpha 1	collagen, t	type XVI,	CCTGGTCCTC [C/T] GGGATTGCCA	υ Σ	E	Д	ы
G712118	WIAF-13863	M92642	3199 a		collagen, t	type XVI,	TCCTGGCTGT [G/T] TTGGGAGCCC	υ Σ	E	٥	[z-i
G712u9	WIAF-13878	M92642	318 8		collagen, t	type XVI,	ACCTCATCCA [C/T] CGACTCAGCC	8	E-1	斑	Ħ
G712u10	WIAF-13882	M92642	1346 a	COL16A1, co alpha 1	collagen, t	type XVI,	ACAGGCGAGA [A/G] GGGCCAGAAA	Æ	_U	×	ద
G712u11	WIAF-13883	M92642	COLLER 1309 alpha	1,	collagen, t	type XVI,	GTCAGGAGCT [C/T] TGGGACCCTC	S	ы	ᆈ	ы
G715a1	WIAF-13344	274615	3504 C	3504 COLIA1, col	collagen, ty	type I, alpha 1 1	TCCTGGTGAA [C/G] AAGGTCCCTC	Σ E	₀	α	Ы
G717u1	WIAF-12639	274616	3988	3988 COL1A2, col	collagen, ty	type I, alpha 2 1	ATGAGGAGAC [T/C] GGCAACCTGA	S	Ü	[+	E
G720u1	WIAF-12367	X14420	3494 a	COL3A1, collagen, 1 (Ehlers-Danlos sy autosomal dominant)	collagen, type -Danlos syndrom dominant)	III, alpha e type IV,	GGTGCAATCG [G/A] CAGTCCAGGA	ڻ ع	A.	U	Д
G720u2	WIAF-12383	X14420	3035	COL3A1, collagen, 1 (Ehlers-Danlos sy autosomal dominant)	lagen, t mlos syn minant)	COL3A1, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant)	GGTGTCAAGG [G/A] TGAAAGTGGG	ت ع	A	ರ	D
G720a3	WIAF-13374	X14420	214 9	COL3A1, collagen, 1 (Ehlers-Danlos sy 214 autosomal dominant)	collagen, type III, -Danlos syndrome tyj dominant)	alpha pe IV,	TCTTGGTCAG [T/C] CCTATGCGGA	E	ט	S	д
G720a4	WIAF-13375	X14420	1953 8	COL3A1, col 1 (Ehlers-De autosomal do	collagen, ty Danlos syndominant)	COL3A1, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant)	CTGGACCTCA [A/G] GGACCCCCAG	8	ত ব	a	Ø
G720a5	WIAF-13376	X14420	2194	COL3Al, col 1 (Ehlers-Da autosomal do	ollagen, t Danlos syn dominant)	COL3A1, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant)	TAGAGGIGGA [G/A] CIGGICCCCC	E	<u>ط</u> ق	Æ	H
G720a6	WIAF-13377	X14420	3731	COL3A1, collagen, 1 (Ehlers-Danlos sy 3731 autosomal dominant)	llagen, t anlos syn ominant)	COL3A1, collagen, type III, alpha 1 (Ehlers-Danlos syndrome type IV, autosomal dominant)	GGGATTGGAG [G/A] TGAAAAAGCT	Σ	<u>ط</u> ق		Ω

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G722u1	WIAF-14132	HT3162	140 2	, 704,	corrager,		r N	GAGATTGGCG[C/T]GACTGGTGAT	Σ	C	ď	Þ
G724a1	WIAF-12120	X81053	3892 4	COL4A4,	collagen,	type IV,	alpha	CTCGTGGAAA [G/A] AAAGGTCCCC	S	G A	×	ೱ
G724a2	WIAF-12121	X81053	4187 4	COL4A4,	collagen,	type IV,	alpha	GAAAGGACCA [A/G] TGGGATTCCC	М	A G	Σ	>
G724a3	WIAF-12122	X81053	3802 4	COL4A4,	collagen,	type IV,	alpha	ATGATGTGGG [G/A] CCACCTGGTC	S	ტ 		U
G724a4	WIAF-12123	X81053	1838 4	COL4A4,	collagen,	type IV,	alpha	accaggaaag[c/a]atggtgcctc	Æ	C B	ш	z
G724u5	WIAF-12364	X81053	376 4	COL4A4,	collagen,	type IV,	alpha	CTGTTTGCCA[C/T]TGTGTTCCTG	S	H ن	- =	н
G724u6	WIAF-12365	X81053	2018 4	COL4A4,	collagen,	type IV,	alpha	TCCAGGGGT [C/G] ATGAAGATGC	E	ق د	H	
G724u7	WIAF-12366	X81053	4756 4	COL4A4, 4	collagen,	type IV,	alpha	GCCTTCCCGT [A/G] TTTAGCACGC	တ	P. G.	>	>
G724u8	WIAF-12377	X81053	3595 4	COL4A4,	collagen,	type IV,	alpha	CTGGACCACC [A/G] GGGTGCCCAG	ω	ڻ لا	Дı	д
G724u9	WIAF-12378	X81053	3516 4	COL4A4,	collagen,	type IV,	alpha	GGAGCATCCG [G/C]AGAGCAGGGC	×	ပ	Ø.	4
G724u10	WIAF-12379	X81053	4288	COL4A4,	collagen,	type IV,	alpha	CTGGTCTTCC [A/G] GGTCCCAGAG	8	₽ G	<u>Д</u>	Д
G724u11	WIAF-12380	X81053	5140 4	COL4A4, 4	collagen,	type IV,	alpha	GCCACTTTT [C/A] GCAAATAAGT	Σ	ر د	Tr.	니
G724u12	WIAF-12387	X81053	207 4	COL4A4, 4	collagen,	type IV,	alpha	GACTTGCCTG [C/T] GATGTGGTCT	1	ا ت		1
G727u1	WIAF-12362	D90279	5135	5135 COL5A1,	collagen,	type V,	alpha 1	TTCAAGGTTT [A/T] CTGCAACTTC	Σ	T.	7	<u>F4</u>
G727u2	WIAF-12369	D90279	4686	4686 COL5A1,	collagen,	type V,	alpha 1	AACAGGGTAT [C/T] ACTGGTCCTT	S	H ت	H	н
G727u3	WIAF-12370	D90279	4608	4608 COL5A1,	collagen,	type V,	alpha 1	TCGGTCCTCC[G/C]GGTGAACAGG	ß	r _O	ρ _ι U	<u></u>
G727a4	WIAF-13300	D90279	2034	2034 COL5A1,	collagen,	type V,	alpha 1	1 ACGGCCTGGC [T/A] GGGTTGCCAG	Ω	T A	A	4
G727a5	WIAF-13301	D90279	2073	2073 COL5A1,	collagen,	type V,	alpha 1	1 GTGACCCTGG [T/C] CCTTCCGGCC	ß	ы	υ U	ט
G727a6	WIAF-13302	D90279	3763	3763 COL5A1,	collagen,	type V,	alpha 1	alpha 1 cGGGCAGAAA[G/A]GTGATGAAGG	Σ	5	A G	S

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G729u1	WIAF-11844	L02870	2345	COLTA1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 2345 recessive)	atggactgga [g/A] ccagalactg	න ආ	4	Д
G729u2	WIAF-11845	102870	3083	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 3083 recessive)	TATCCTGGCG [G/A] CCACTCAGAG	ى ق	A	K K
G729u3	WIAF-11846	1.02870	3031	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 3031 recessive)	GACTCGGTGA [C/T] TTTGGCCTGG	ں بع	FI	H
G729u4	WIAF-11851	102870	1289	COL7Al, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1289 recessive)	CGGACTATGA [G/T] GTGACCGTGA	<u>დ</u>	<u>H</u>	전 다
G729u5	WIAF-11852	L02870	1032	COL7Al, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1032 recessive)	CCAAGTGACT [G/T] TGATTGCCCT	<u>უ</u>	Eн	
G729u6	WIAF-11853	L02870	1897	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1897 recessive)	CGCCGGGAGC [C/T] GGAAACTCCA	<u>ე</u>	٤٠	<u>ы</u>
G729u7	WIAF-11854	L02870	1827	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1827 recessive)	GCTTAGCTAC [A/T] CTGTGCGGGT	<b>E</b>	E	Ε- (S
G729u8	WIAF-11855	L02870	1893	COLTAL, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1893 recessive)	TGTCCGCCGG [G/A] AGCCGGAAAC	<u>უ</u>	A	M M

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G729u9	WIAF-11864	102870	2142	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 2142 recessive)	GGGCCCTGCT [G/A] CAGTCATCGT	M D		F
G729u10	WIAF-11865	L02870	2353	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 2353 recessive)	GAGCCAGATA [C/T] TGAGTATACG	E E	H	Н
G729u11	WIAF-11866	L02870	2221	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 2221 recessive)	TCATCTGTCA [C/T] CATTACCTGG	E4 U E	E	н
G729u12	WIAF-11869	102870	6583	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 6585 recessive)	ACCAGGAGAG [C/T] GTGGTATGGC	E E	<u> </u>	υ
G729u13	WIAF-11870	102870	8169	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 8169 recessive)	GGGTGACCGA [G/T] GCTTTGACGG	ნ- ტ დ	<b>o</b>	υ
G729u14	WIAF-11877	L02870	4. 8. 8.	COL/7Al, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 438 recessive)	CGCCATCCGT [G/A] AGCTTAGCTA	ط ن ک	লে	×
G729u15	WIAF-11882	L02870	3481	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 3481 recessive)	AGGATCCGTG [A/T] CATGCCCTAC	М 4	Ð	Δ
G729u16	WIAF-11883	L02870	5654	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 5654 recessive)	ACGGAGAACC [T/C] GGGGACCCTG	ω Η		Д

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G729u17	WIAF-11884	102870	7124	COLTA1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 7124 recessive)	TGCCAGGGC [G/C] CGAGGCGAGA	<u>ა</u>	υ	д	<u>С</u> ч
G729u18	WIAF-11885	102870	7277	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 7757 recessive)	GCTTGGATGG [T/C] GACAAAGGAC	<u>ا</u>	Ü	Ü	ט
G729u19	WIAF-13389	L02870	1615	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 1615 recessive)	ACCGTGGTTC [C/T] CACTGGACCA	<u>ن</u> ع	E-1	വ	L1
G729u20	WIAF-13390	L02870	2930	COL7Al, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 2930 recessive)	TCCTAGGGCC [G/A]GCTGGAGAAG	ر م	4	Д	Ωι
G729u21	WIAF-13399	L02870	5145	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 5145 recessive)	CCAGGGAGAT [C/T] CTGGAGAGGA	υ  Σ	E	ρι	w
27.00.1.00	WT2F-12411	1.02870	3472	COL7A1, collagen, type VII, alpha 1 (epidermolysis bullosa, dystrophic, dominant and 3472 recessive)	ATCTTGCAAA [G/A] GATCCGTGAC	ъ Б	A.	ద	×
G730a1	WIAF-13303	X57527	305	COL8A1, collagen, type VIII,	ATGGGCAAGG [A/G] AGCCGTTCCC	Æ	ڻ ا	网	ro l
G732u1	WIAF-12616	M95610	936 2	COL9A2, collagen, type IX, alpha	CAGGCGGGAC [A/G] GCCCGGAAGT	S A	ರ	E	Ę÷
G732u2	WIAF-12617	M95610	696 2	COL9A2, collagen, type IX, alpha 2	AAGGGAGAGA [C/T] GGGCCCTCAT	ა ა	E⊣	Д	Д
G732u3	WIAF-12619	M95610	1288 2	COL9A2, collagen, type IX, alpha	AAGTGGGTGA [C/T] CCAGGGGTGG	υ Σ	E→	д	S
G732u4	WIAF-12620	M95610	962 2	COL9A2, collagen, type IX, alpha 2	CCACCAGGG [C/G] TAGCGGGTGT	υ Σ	Ŋ		ద

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WIAF-13394	M1343	٥.	ct A	TGCTCCCTG [G/L]	٠.			
WIAF -13383	M5854	183		ATGGAGGCT [A/G] AAGTCCAAGA	E	A G	×	田
4F-13384	M58549	330	330 MGP, matrix Gla protein	GCGCCGAGGG [A/G] CCAAATGAGA	M	A G	H	Ą
WIAF-11867	U94332	862	TNFRSF11B, tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin)	TGCTGAAGTT [A/G] TGGAAACATC	S	A 0		
WIAF-11874	U94332	1244	TWFRSF11B, tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin)	GTATCAGAAG [7/C] TATTTTTAGA	0.0			<u> </u>
WIAF-13402	HT847	1669	PTHR1, parathyroid hormone receptor 1	CCCTGGAGAC [C/A] CTCGAGACCA				1 1
WIAF-12414	J03040	123	SPARC, secreted protein, acidic, cysteine-rich (osteonectin)	CTCAGCAAGA [A/G] GCCCTGCTG				<u> </u>
WIAF-12628	HT0157	117	VDR, vitamin D (1,25-dihydroxyvitamin D3) receptor	CCTTCAGGGA [T/C] GGAGGCAATG	Σ	D E	Σ	E
WIAF-12629	HT0157	1171	VDR, vitamin D (1,25- 1171 dihydroxyvitamin D3) receptor	CCGCGCTGAT [T/C] GAGGCCATCC	ω _.	L U	Н	н
WIAF-12640	HT0157	172	VDR, vitamin D (1,25- 172 dihydroxyvitamin D3) receptor	TTGACCGGAA [C/T]GTGCCCGGA	Ω.	U U	Z	z
WIAF-11862	HT3734	679	679 osteopontin, alt. transcript 1	ATCACCTCAC [A/T] CATGGAAAGC	M A	A	田田	1
WIAF-11875	HT3734	386	386 osteopontin, alt. transcript 1	AAGATGATGA [A/G] GACCATGTGG	S	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	Б	Д
WIAF-11876	HT3734	419	419 osteopontin, alt. transcript 1	CCATTGACTC [G/A] AACGACTCTG	ω O	ى د	တ	တ
WIAF-12084	HT3734	171	osteopontin, alt. transcript 1	TAAACAGGCT [G/A]ATTCTGGAAG	Σ		Д	Z
WIAF-13387	HT3734	738	osteopontin, alt. transcript 1	CCAGGACCTG [A/C] ACGCGCCTTC	M A	A C	Z	н
WIAF-13388	HT3734	716	g	CATACAAGGC [C/A] ATCCCCGTTG	S	4	_ ≪	A
W1AF-12631	HT5036	410	410 ADM, adrenomedullin	ひつびつつびかるひつ [ひ/ひ] ひ上むないなないなひ	5	7	,	D

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G752u1	WIAF-11843	HT1782	CHGA, chromogranin A (parathyroid 1405 secretory protein 1)	oid CGGCCATTGA[A/G]GCAGAGCTGG	8 4	Ŋ	[22]	<u>F</u>
G752u2	WIAF-11873	HT1782	CHGA, chromogranin A (parathyroid 1187 secretory protein 1)	oid GGACAACCGG[G/A]ACAGTTCCAT	<u>ت</u>	Æ	Д	z
G754a1	WIAF-13382	K02043	NPPA, natriuretic peptide 663 precursor A	GTACAATGCC [G/A] TGTCCAACGC	∑ ∑	Ą	>	Σ
G756u1	WIAF-12395	HT3508	SCNN1A, sodium channel, 2086 nonvoltage-gated 1 alpha	CAGTTCCTCC [A/G] CCTGTCCTCT	M	r _O	E	Æ
G757u1	WIAF-12420	HT28563	SCNNIB, sodium channel, nonvoltage-gated 1, beta (Liddle 797 syndrome)	CCTGCAGGCC [A/C] CCAACATCTT	<u>Σ</u>	U	H	Д
G757u2	WIAF-12421	HT28563	SCNN1B, sodium channel, nonvoltage-gated 1, beta (Liddle 1006 syndrome)	GAACTGAATT [C/T] GGCCTGAAGT	<u>v</u>	H	ഥ	Ŀ
G757u3	WIAF-12430	HT28563	SCNNIB, sodium channel, nonvoltage-gated 1, beta (Liddle 1768 syndrome)	TCATCGACTT [T/C] GTGTGGATCA	S E	ט	Ţ.	ţri
G757u4	WIAF-12494	HT28563	SCNN1B, sodium channel, nonvoltage-gated 1, beta (Liddle 662 syndrome)	AAGCAGCTCA [G/C] CATCAGAAAA	<u></u> છ	ū	Ą	Ωŧ
G757u5	WIAF-12506	HT28563	SCNNIB, sodium channel, nonvoltage-gated 1, beta (Liddle 1091 syndrome)	e GATGCTTCAC [G/C] AGCAGAGGTC	Œ	ט	<u> </u>	α
G757u6	WIAF-12507	HT28563	SCNNIB, sodium channel, nonvoltage-gated 1, beta (Liddle 1452 syndrome)	ACCIGCATIG [G/I] CALGIGCAAG	υ Σ	H	ರ	Þ
G758u1	WIAF-12621	HT27856	SCNNID, sodium channel,	CGGGAACCCA [C/T] GTCGGCCGAG	Σ Σ	H	£	ŭ
G758u2	WIAF-12632	HT27856	SCNNID, sodium channel, 325 nonvoltage-gated 1, delta	CCTCTTTGAG [C/T] GTCACTGGCA	U ∑	H	ద	U
G758u3	WIAF-12634	HT27856	SCNNID, sodium channel, 879 nonvoltage-gated 1, delta	ATGGCGTCTG [G/A] ACAGCTCAGC	D N	Æ	- 3	*
G758u4	WIAF-12635	HT27856	SCNNID, sodium channel, 1138 nonvoltage-gated 1, delta	CGTGGAGGTG [G/C] AGCTGCTACA	Æ	Ü	ম	Ø
G762u1	WIAF-12622	HT27531	NPR3, natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor	or TAGGAGCTGG[C/T]TTGCTAATGG	υ υ	E+	ტ	_ ღ

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G762u2	WIAF-12623	HT27531	NF re (a) (a) (1926 C)	R3, natriuretic peptide ceptor C/guanylate cyclase C trionatriuretic peptide receptor	AGAAGAAAGT [A/G]ACCTTGGAAA	M	ರ	z	Д
G762u3	WIAF-12624	HT27531	NF re (a 1791 C)	R3, natriuretic peptide ceptor C/guanylate cyclase C trionatriuretic peptide receptor	CAAATCATCA [G/T] GTGGCCTAGA	B M	EH:	უ	ن
G762u4	WIAF-12636	HT27531	NF re (a 1963 C)	R3, natriuretic peptide ceptor C/guanylate cyclase C trionatriuretic peptide receptor	GAAGATTCCA [T/C] CAGATCCCAT	는 돈	ن	н	H
G763u1	WIAF-12659	HT3183	NF re (a) 1633 B)	R2, natriuretic peptide ceptor B/guanylate cyclase B trionatriuretic peptide receptor	CTGGGCCCTT [C/T] CCTGATGAAC	<u>Σ</u>	H	Ø	ΪΞŧ
G763u2	WIAF-12678	HT3183	NP re (a (a 668 B)	R2, natriuretic peptide ceptor B/guanylate cyclase B trionatriuretic peptide receptor	TGCCATCACT [T/C] CTGCTGTTGG	S	υ		Ţ
G763u3	 WIAF-12684	HT3183	NF re (a 2354 B)	R2, natriuretic peptide ceptor B/guanylate cyclase B trionatriuretic peptide receptor	TGTTTGAACT [C/T] AAACATATGA	ນ s	F	ц	딘
G764u1	WIAF-12698	HT1221	3021	NPR1, natriuretic peptide receptor A/guanylate cyclase A (atrionatriuretic peptide receptor A)	CCCGGTTACT [G/T] ICTCTTTGGG	<u>უ</u>	E-1	ن	[14
G764u2	WIAF-12708	HT1221	NH re (2 (8	RI, natriuretic peptide sceptor A/guanylate cyclase A itrionatriuretic peptide receptor	GAGCGCCAAG [C/T] GCTCATGCTC	υ Σ	E-i	4	<b>&gt;</b>

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			A C	NPR1, natriuretic peptide receptor A/guanylate cyclase A				
G764u3	WIAF-12709	HT1221	1897 A)	utrionatriuretic peptide receptor	GTCCCCGTGG [G/A] AGCCTGCAGG	თ დ	4	D D
G765u1	WIAF-10012	HT2456	604	DCP1, dipeptidyl carboxypeptidase 1 (angiotensin I converting 604 enzyme)	GCTGGCACAA [A/G] GCTGCGGGCA	ଅ ୟ	ტ	z z
G765u2	WIAF-10014	HT2456	2350	DCP1, dipeptidyl carboxypeptidase 1 (angiotensin I converting	TGATGGCCAC [A/G] TCCCGGAAAT	ر ا		En En
G765u3	WIAF-10025	HT2456	1688	DCP1, dipeptidyl carboxypeptidase 1 (angiotensin I converting 1688 enzyme)	CCCACTGCAC [C/A] AGTGTGACAT	ت ع	≮	о Я
G765u4	WIAF-10027	HT2456	3220	DCP1, dipeptidyl carboxypeptidase 1 (angiotensin I converting 3220 enzyme)	TCCCCTTCAG [C/T] TACCTCGTCG	ე გ	EH.	S S
G765u5	WIAF-10028	HT2456	3409	DCP1, dipeptidyl carboxypeptidase 1 (angiotensin I converting 3409 enzyme)	TCAGGTACTT [T/C] GTCAGCTTCA	Ω <u></u> ⊟	۲	Er Er
G765u6	WIAF-10040	HT2456	775	dipeptidyl carboxypeptidase otensin I converting	AGCCCCTCTA [C/T] CTGAACCTCC	ა ე	E	X X
G772u1	WIAF-12626	HT2121	1064	AVPR2, arginine vasopressin receptor 2 (nephrogenic diabetes 1064 insipidus)	TCAGCAGCAG [C/T] GTGTCCTCAG	<u>ი</u>		S S
G772u2	WIAF-12627	HT2121	866	AVPR2, arginine vasopressin receptor 2 (nephrogenic diabetes insipidus)	CCTITGIGCT [A/G] CICATGITGC	رم الا	೮	<u>П</u>
G773u1	WIAF-12644	HT2141	163	SLC6A6, solute carrier family 6 (neurotransmitter transporter, 163 taurine), member 6	CTAGCAAGAT [C/T] GACTTTGTGC	_ي م	F	Н

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G773u2	WIAF-12645	HT2141	445	SLC6A6, solute carrier family 6 (neurotransmitter transporter, 445 taurine), member 6	TCGTCATCCT [G/C] GCCTGGGCCA	დ დ	υ	н	н
G773u3	WIAF-12665	HT2141	289	SLC6A6, solute carrier family 6 (neurotransmitter transporter, 289 taurine), member 6	TGTTTGGGAG [C/T] GGCCTGCCTG	න ව	H	Ŋ	ß
G773u4	WIAF-12666	HT2141	382	SLC6A6, solute carrier family 6 (neurotransmitter transporter, 382 taurine), member 6	CCTIGITCTC [I/C] GGTATCGGCT	Ω ⊟	ن	S	တ
G776u1	WIAF-11857	U66088	1457	SLC5A5, solute carrier family 5 (sodium iodide symporter), member 5	TAGAAGACCT[C/T]ATCAAACCTC	ე ე	FI	ı,	L
G776u2	WIAF-11871	066088	2039	SLC5A5, solute carrier family 5 (sodium iodide symporter), member 5	GATTGTTGTG [G/C] TGGGACCTCG	<u>დ</u>	υ	×	υ
G776u3	WIAF-13398	066088	1379	SLC5A5, solute carrier family 5 (sodium iodide symporter), member 5	GGCTTTTCCT [G/A] GCCTGTGCTT	ය	Ą	П	ы
G777u1	WIAF-12646	HT27843	4348 SMRT	SMRT	ATACAATATC [A/G] GCCAGCCTGG	MA	ß	လ	ro O
G777u2	WIAF-12654	HT27843	2031 SMRT	SMRT	CTGAGCTGGG [T/C] AAGCCGCGGC	S3 E4	ပ	IJ	ט
G777u3	WIAF-12655	HT27843	2052 SMRT	SMRT	AGAGCCCCCT [G/A] ACCTATGAGG	S G	Ą	П	П
G777u4	WIAF-12675	HT27843	2205 SMRT	SMRT	CTCGTGAGAT [C/T] GCCAAGTCCC	S C	ㅂ	н	н
G778u1	WIAF-14093	HT1449	8212 TG,	TG, thyroglobulin	ATCTCGTCTC [T/C] GAAGACATCT	M	ပ	Д	Д
G778u2	WIAF-14111	HT1449	6033 TG,	TG, thyroglobulin	ATGTGAACGA [C/T] GGTGCGATGC	N C	Ħ	ద	М
G778u3	WIAF-14112	HT1449	6894 TG,	TG, thyroglobulin	GTATCTCAAT [G/T] TGTTCATCCC	MG	Т	Λ	Ţ
G778u4	WIAF-14125	HT1449	2375 TG,	TG, thyroglobulin	ATGGGCCTCC [T/C] GAGCAGGTCT	S	ပ	д	д
G778u5	WIAF-14136	HT1449	1931 TG,	TG, thyroglobulin	AGGATGTCCA [A/G] TGCTTTTCCG	S	D	α	a
G783u1	WIAF-12649	X97674	4008	H.sapiens mRNA for transcriptional 4008 intermediary factor 2.	CTAGTGGTAT [G/C] CCAGCAACTA	M G	Ü	Σ	н
G783u2	WIAF-12658	X97674	2566	H.sapiens mRNA for transcriptional 2566 intermediary factor 2.	GCCTGGCAGT [G/A] AGCTGGACAA	<u>უ</u>	Æ	臼	×
G783u3	WIAF-12671	X97674	3828	H.sapiens mRNA for transcriptional 3828 intermediary factor 2.	CTCTGAGGCC [T/C] GGAGTACCAA	S H	U	д	Д
G785u1	WIAF-13385	HT1291	386	TTR, transthyretin (prealbumin, 386 amyloidosis type I)	CCAACGACTC [C/T] GGCCCCGCC	S D	E	Ø	S

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G787u1	WIAF-12652	HT27477	468	TRIP15: thyroid receptor 468 interacting protein 15	GAAAATTATA [T/C] TTAGAACGAG	S F	ນ	≯	ÞI
G792u1	WIAF-12661	HT27476	265	thyroid receptor interactor 14	CAGCTGGAAC [G/A] TGAAGAGGGC	_ Σ	ď	Þ	Σ
G793u1	WIAF-12643	HT5152	458	thyroid receptor interactor 8	GGAAGCTTTT[C/G]AAAGAATGTT	U Z	හ	ഗ	*
G794u1	WIAF-12664	HT5136	1110	PSMC5, proteasome (prosome, 1110 macropain) 26S subunit, ATPase, 5	GCGTGTGCAC [G/A] GAAGCTGGCA	დ	A.	EH	[-
G797u1	WIAF-11847	HT3919	140	140 glutamate receptor 3, flip isoform	isoform CTCACGGAGG[A/G]TTCCCCAACA	S	_O	r r	ט
G797u2	WIAF-11848	HT3919	759	glutamate receptor 3, flip	isoform GGTTGTGATC[C/T]TAGGGAAACA	ა ა	E	ᄓ	ы
G797u3	WIAF-11849	HT3919	1253	glutamate receptor 3, flip	isoform GCTACTGGAA[C/T]GAGTATGAAA	ပ ပ	E⊣	z	z
G797u4	WIAF-11850	HT3919	1770	1770 glutamate receptor 3, flip isoform	isoform TCTTTTCCTA[G/A]TCAGCAGGTT	υ Σ	Æ	>	н
G797u5	WIAF-13404	HT3919	2711	2711 glutamate receptor 3, flip isoform	isoform GCTACAACGT[G/A]TATGGAACAG	S	4,	>	٥
G797u6	WIAF-13405	HT3919	2376	2376 glutamate receptor 3, flip isoform	isoform CTCAGCATTA[G/A]GAACGCCTGT	<u>უ</u>	Ø	ڻ ت	딵
G798u1	WIAF-11868	X77748	2655	GRM3, glutamate receptor, 2655 metabotropic 3	TGCAGACGAC [A/G] ACCATGTGCA	ج د	ŋ	EH	EH
G798u2	WIAF-11879	X77748	2771	GRM3, glutamate receptor, 2771 metabotropic 3	CACAGACTGC [A/G] CCTCAACAGG	A A	U	耳	DK.
G798a3	WIAF-12085	X77748	2699	GRM3, glutamate receptor, 2699 metabotropic 3	GIGGICTIGG [G/C] CIGITIGITI	<u>υ</u>	Ü	ט	A
G798a4	WIAF-12086	X77748	2738	GRM3, glutamate receptor, 2738 metabotropic 3	ATCCTGTTTC [A/G] ACCCCAGAAG	MA	ტ	α	ద
G798a5	WIAF-12087	X77748	2072	GRM3, glutamate receptor, 2072 metabotropic 3	ACACCCTTGG [T/C] CAAAGCATCG	Σ F	υ	٥	ď
G798a6	WIAF-12088	X77748	2235	GRM3, glutamate receptor, 2235 metabotropic 3	CCCTGCTGAC[C/T]AAGACAAACT	ე ე	Ę-4	E⊣	ы
G798u7	WIAF-13391	X77748	1131	GRM3, glutamate receptor, 1131 metabotropic 3	GCGCCAATGC [C/T] TCCTTCACCT	ນ ນ	E	Æ	ď
G799u1	WIAF-11880	M81883	2000	GAD1, glutamate decarboxylase 1 2000 (brain, 67kD)	CAACAAATGC[C/T]TGGAACTGGC	S C	H	н	ı
G799u2	WIAF-11881	M81883	1822	GAD1, glutamate decarboxylase 1 1822 (brain, 67kD)	AGGGTATACT [C/T] CAAGGATGCA		H		н

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G799u3	WIAF-13392	M81883	661	GAD1, glutamate decarboxylase 1 (61 (brain, 67kD)	GCGTGGCCCA[T/C]GGATGCACCA	S	ບ	н	Н
G799u4	WIAF-13393	M81883	556	GAD1, glutamate decarboxylase 1 556 (brain, 67kD)	AGCTGATGGC [G/A] TCTTCGACCC	ა ზ	A	A.	A
G799u5	WIAF-13410	M81883	1229	GAD1, glutamate decarboxylase 1 (brain, 67kD)	CCTCATGGAA [C/T] AAATAACACT	N N	E+	α	*
G801u1	WIAF-13403	D49394	1596	HTR3, 5-hydroxytryptamine (serotonin) receptor 3	TTTACCTGCT [A/G] GCGGTGCTGG	S A	ŋ	1	ı
G803a1	WIAF-13118	U66406	1446	1446 EFNB3, ephrin-B3	CTGGGCCTGG [G/A] GGGTGGAGGT	Σ U	Æ	Ð	凶
G804u1	WIAF-11887	Z26653	7237	LAMA2, laminin, alpha 2 (merosin, 7237 congenital muscular dystrophy)	1, TCACTGATGG [G/T] CACATAAAAG	ა ე	E⊣	<u> </u>	
G804u2	WIAF-11901	726653	9351	LAMA2, laminin, alpha 2 (merosin, congenital muscular dystrophy)	a, GCAAGCCACT[G/C]GAGGTTAATT	υ Σ	υ	Z	ω
G804u3	WIAF-11924	Z26653	8740	LAMA2, laminin, alpha 2 (merosin, 8740 congenital muscular dystrophy)	a, ACACTACCCG [A/G] AGAATTGGTC	S	<u>.</u>	ద	<b>R</b>
G804u4	WIAF-11943	226653	8577	LAMA2, laminin, alpha 2 (merosin, 8577 congenital muscular dystrophy)	n, ACCAAAATCA[A/G]TGATGGCCAG	M	ט	Z	S
G804a5	WIAF-12089	Z26653	3372	LAMA2, laminin, alpha 2 (merosin, congenital muscular dystrophy)	n, crongraer [6/a] crrccrocr	M	Æ	۲	K
G804a6	WIAF-13227	226653	7047	LAMA2, laminin, alpha 2 (merosin, 7047 congenital muscular dystrophy)	n, GTCAGTCCTC[A/g]GGTGGAAGAT	M	<u>σ</u>	0	ద
G804u7	WIAF-13437	226653	6791	LAMA2, laminin, alpha 2 (merosin, 6791 congenital muscular dystrophy)	n, TGTGAGAGCC[C/T]TGGATGGACC	ა ა	<u></u>	FI	Ы
G805u1	WIAF-13416	U14755	799	799 LHX1, LIM homeobox protein 1	AAGTAACAGC [A/G] GTGTTGCCAA	M	rg d	ശ	Ö
G805u2	WIAF-13417	U14755	743	743 LHX1, LIM homeobox protein 1	GGCGAGGAAC [T/C] CTACATCATC	×	υ	긔	
G805u3	WIAF-13428	U14755	639	639 LHX1, LIM homeobox protein 1	GCCGTCAGGG [C/A] ATCTCCCCTA	ى د	4	Ü	_o
G806u1	WIAF-11886	AF026547	2656	CSPG3, chondroitin sulfate 2656 proteoglycan 3 (neurocan)	TTGGAGTTCC [A/G] GCCATGTCTA	8	<u> </u>	Сц	<u>C</u> .

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G806u2	WIAF-11895	AF026547	CS 529 pr	CSPG3, chondroitin sulfate 529 proteoglycan 3 (neurocan)	TGACCTTCGC [T/C] GAGGCCCAGG	S Ed	υ	Ą	Æ
G806u3	WIAF-11896	AF026547	CS 477 pr	CSPG3, chondroitin sulfate proteoglycan 3 (neurocan)	GAGGTGACAG [G/A] TGTTGTGTTC	<u>უ</u>	Æ	ტ	Д
G806u4	WIAF-11917	AF026547	CS 89 pr	CSPG3, chondroitin sulfate 89 proteoglycan 3 (neurocan)	ACAGGATATC [A/G] CCGATGCCAG	Σ	ט	E	A.
G806u5	WIAF-11918	AF026547	CS 213 pr	CSPG3, chondroitin sulfate 213 proteoglycan 3 (neurocan)	AGGGCAGCCC [G/C] AGATGCCCCT	. හ න	U	D4	д
980ene	WIAF-11929	AF026547	CS 769 pr	CSPG3, chondroitin sulfate 769 proteoglycan 3 (neurocan)	GCTTTGCCCG [G/A] GAGCTGGGGG	ა დ	Ą	p4	PK.
G806u7	WIAF-11931	AF026547	CS 3148 pr	CSPG3, chondroitin sulfate 3148 proteoglycan 3 (neurocan)	ACATTGATGA [C/T] TGCCTCTGCA	ပ အ	H	А	Д
80908£	WIRF-11949	AF026547	209 pr	CSPG3, chondroitin sulfate 209 proteoglycan 3 (neurocan)	GCCAAGCGCA [G/A] CCCGAGATGC	<u>ტ</u>	Æ	A	E
G806a9	WIAF-13114	AF026547	3430 pr	CSPG3, chondroitin sulfate 3430 proteoglycan 3 (neurocan)	ATGAAAACAC [G/A] TGGATCGGCC	ა ე	A	EH	E
G806u10	WIAF-13420	AF026547	2113 pr	CSPG3, chondroitin sulfate 2113 proteoglycan 3 (neurocan)	CCAGGGCAGA [C/G] TTCAGAGAAA	υ Σ	. U	О	臼
G806u11	WIAF-13431	AF026547	94 pr	CSPG3, chondroitin sulfate 94 proteoglycan 3 (neurocan)	ATATCACCGA [T/G] GCCAGCGAAA	E E	უ	Д	EXI
G806u12	WIAF-13432	AF026547	CE 275 pr	CSPG3, chondroitin sulfate 275 proteoglycan 3 (neurocan)	ACAGGACTTG [C/T] CCATCCTGGT	υ Σ	E	O.	ß
G808a1	WIAF-13117	Y13276	IT (I	TLX, tailless homolog (Drosophila)	GCATGAGCAA [G/a] CCAGCCGGAT	S	ಹ	×	×
G810u1	WIAF-11890	X98248	990 SORT1,	ORT1, sortilin 1		S C	Æ	E	E⊣
G810u2	WIAF-11891	X98248	1093 SORT1,	ORT1, sortilin 1	GGCAGCAAAT [G/T] ATGACATGGT	<u>υ</u>	Н	Д	¥
G810u3	WIAF-11907	X98248	1683 SORT1,	ORT1, sortilin 1	CAGACGAAGG [T/G] CAATGCTGGC	S	ט	- 1	Ü
G810u4	WIAF-11908	X98248	1433 SORT1,	ORT1, sortilin 1		Æ	υ		드
G810u5	WIAF-11909	X98248	1354 SORT1,				O .		
G810u6	WIAF-11910	X98248	2180 SORT1,	sortilin		E I C	K K	<b>н</b> с	z z
G810u7	WIAF-11911	X98248	2264 SORII,	ORTI, sortilin 1	AACITITIGA [6/A] ICCGGAAAAA	1	4	0	4

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G810u8	WIAF-11925	X98248	1993	1993 SORT1, sortilin 1	TCGAGACTAT [G/A] TTGTGACCAA	M G A V	н
G810u9	WIAF-11939	X98248	1351	1351 SORT1, sortilin 1	GAGGAAGCCT [G/C] AAAACAGTGA M	M O O	ø
G810u10	WIAF-11940	X98248	2232	SORT1, sortilin 1	AAGTAAAAGA [C/T] TTGAAAAAGA	E D	
G810a11	WIAF-13115	X98248	1769	SORT1, sortilin 1	TCCATGAATA [T/A] CAGCATTTGG M	T A	
G810a12	WIAF-13116	X98248	1757	1757 SORT1, sortilin 1	CCTGGAGCTA [G/A] GTCCATGAAT M	M G A	×
G811u1	WIAF-11893	HT3676	3 006	900 synapsin I, alt. transcript 1	TGACCAAGAC [G/A] TATGCCACTG	S S	E
G811u2	WIAF-11894	HT3676	758 8	758 synapsin I, alt. transcript 1	ACCTTCTACC[C/T]CAATCACAAA	M C H	ㅁ
G811u3	WIAF-11927	HT3676	966	synapsin I, alt. transcript 1	CGTCAGTGTC [A/T] GGGAACTGGA	S T S	လ
G811u4	WIAF-11928	HT3676	1054 8	synapsin I, alt. transcript 1	CATGTCTGAC[A/G]GATACAAGCT M	Σ Δ	ט
G811u5	WIAF-13418	HT3676	249	synapsin I, alt. transcript 1	TGTCCAACGC [G/A] GTCAAGCAGA	S G A	Æ
G811u6	WIAF-13419	HT3676	432	synapsin I, alt. transcript 1	TTAAAGTAGA [G/A] CAGGCCGAAT	ম ম ম	<u>M</u>
G812n1	WIAF-11898	HT4564	163	163 STX1A, syntaxin 1A (brain)	CCAACCCCGA [T/C]GAGAAGACGA	S T C D	_Д
G812u2	WIAF-11942	HT4564	604	STX1A, syntaxin 1A (brain)	TACACGACAT [G/T] TTCATGGACA	E E	H
G813u1	WIAF-11934	U72508	939	Human B7 mRNA, complete cds.	TATGACAGAG [G/A] ACAGAGGATG	Μ Θ Α	E
G813u2	WIAF-11948	U72508	619	Human B7 mRNA, complete cds.	GCATCCACAT [G/C] GTGACAGGTC	M G M	Н
G816u1	WIAF-11897	HT4230	151	HTR2B, 5-hydroxytryptamine (serotonin) receptor 2B	CTAACTGGTC [T/G] GGATTACAGA	හ E1	w w
G816u2	WIAF-11930	HT4230	189	HTR2B, 5-hydroxytryptamine 189 (serotonin) receptor 2B	GAAATGAAC [A/G] GATTGTTGAG	۳ م	<u>ж</u> О
G818u1	WIAF-11902	HT2694	753	TPH, tryptophan hydroxylase (tryptophan 5-monooxygenase)	GAGTTTTTCA [C/T] TGCACTCAAT	υ ω	н
G818u2	WIAF-11903	HT2694	775	TPH, tryptophan hydroxylase (tryptophan 5-monooxygenase)	TGTGAGACAC [A/G] GTTCAGATCC	о Д Ж	o o
G818u3	WIAF-11904	HT2694	1211	TPH, tryptophan hydroxylase (tryptophan 5-monooxygenase)	TATAATCCAT [A/C] TACACGGAGT	υ «	Ω

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G818u4	WIAF-11905	HT2694	1081	TPH, tryptophan hydroxylase (tryptophan 5-monooxygenase)	GATTACCTGC [A/C] AACAGGAATG	M	υ	M	Ø
n01	WT # 11 19 2 2	HT2694	795		CCTICTATAC [C/T] CCAGAGCCAG	ა ა	E	H	E⊣
91.81	WTAF-11935	HT2694	1239		TCCTGAAAGA [C/T]ACCAAGAGCA	ນ ຮ	E	Д	А
G822u1	WIAF-11906	HT0207	9361		CAGACGGAAA [G/T] TGCTCACACC	<u>დ</u>	F	×	z
G822u2	WIAF-11919	HT0207	637	ASMT, acetylserotonin N-637 methyltransferase	TGGTGGGACA [C/T] GGATAAAGCT	υ Σ	E	ద	×
G822u3	WIAF-11936	HT0207	318	ASWT, acetylserotonin N-318 methyltransferase	GAAAAGCTTT [C/T] TATCGAAACA	ა ე	E	<u>[14</u>	[±,
G822u4	WIAF-11937	HT0207	116	ASMT, acetylserotonin N- 116 methyltransferase	AATGACTACG [C/T] CAACGGCTTC	U E	E-I	A	D
G822u5	WIAF-11938	HT0207	930	ASMT, acetylserotonin N- methyltransferase	ACTGGGCAGA [C/T] GGAAAGTGCT	ပ ပ	E	Δ	Д
G822u6	WIAF-13427	HT0207	120	ASMT, acetylserotonin N- 120 methyltransferase	ACTACGCCAA [C/A]GGCTTCATGG	υ Σ	Æ	Z	×
280 Firl	WTAF-11888	HT4974	236	ADAR, adenosine deaminase, RNA-specific	GCTCAGATAC [C/T] AGCAGCCTGG	U Z	E	Ø	*
1 C C C C C C C C C C C C C C C C C C C	MTAF-11900	HT4974	3076	ADAR, adenosine deaminase, RNA-3076 specific	TCTTTGACAA [A/G] TCCTGCAGCG	Ω <b>4</b>	ტ	_ ⊭	×
2000	WTAR-11912	HT4974	2537	ADAR, adenosine deaminase, RNA-specific	CTTCATTGGG [G/C] AGAACGAGAA	Σ U	Ü	闰	α
28.25.00 28.25.00	WTAF-11941	HT4974	3558	ADAR, adenosine deaminase, RNA-558 specific	GATGGCTATG [A/G] CCTGGAGATC	Æ Æ	ß	Д	ტ
20 CC CC CC CC CC CC CC CC CC CC CC CC CC	WIAF-12090	HT4974	1305	ADAR, adenosine deaminase, RNA-1305 specific	CCTGAGACCA [A/G] AAGAAACGCA	M	ರ	×	ద
G825u6	WIAF-13426	HT4974	3683	ADAR, adenosine deaminase, RNA-specific	CCGCAGGGAT [C/T] TACTGAGACT	S	H	ᆈ	니
G826u1	WIAF-12554	X99383	2109	ADARB1, adenosine deaminase, RNA- 2109 specific, B1 (homolog of rat RED1)	denosine deaminase, RNA- B1 (homolog of rat RED1) AGATTACCAA[A/G]CCCAACGTGT	8	ט	×	×
G826u2	WIAF-12566	X99383	1698	ADARB1, adenosine deaminase, specific, B1 (homolog of rat	RED1) TGTCCTGCAG[T/G]GACAAGATTG	E E	rg l	w	以
G829u1	WIAF-13735	U49262	1404	DVL3, dishevelled 3 (homologous	GGGTTGGAGG [T/C] CCGTGACTGC	Ε	۲	>	A

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G83u1	WIAF-10449	HT1576	1338 m	DNMT1, DNA (cytosine-5-)-1338 methyltransferase 1	ATGATGACCC [G/A] TCTCTTGAAG	ъ В	д	Д
G83u2	WIAF-10450	HT1576	1871 m	DNMT1, DNA (cytosine-5-)- 1871 methyltransferase 1	AAGCTGGTCT [A/G] CCAGATCTTC M	ъ В	Ж	ŭ
G83u3	WIAF-10468	HT1576	928 m	DNMT1, DNA (cytosine-5-)- 928 methyltransferase 1	AAATCCACAG [A/G] TTTCTGATGA M	۵ 9	Н	٥
G83u4	WIAF-10469	HT1576	1562 п	DNMT1, DNA (cytosine-5-)- 1562 methyltransferase 1	AATTCCGACT [C/T] GACCTATGAG M	H ن	တ	Ľ
G83u5	WIAF-10471	HT1576	2424 m	DNMT1, DNA (cytosine-5-)- methyltransferase 1	GGGCCACGTC [G/A] GACCCTCTGG	G A	လ	တ
G83u6	WIAF-10473	HT1576	3790 m	DNWT1, DNA (cytosine-5-)- 3790 methyltransferase 1	GTTCTTCCTC [C/T] TGGAGAATGT S	E U	ы	д
G83u7	WIAF-10486	HT1576	1581	DNMT1, DNA (cytosine-5-)- 1581 methyltransferase 1	AGGACCTGAT [C/A] AACAAGATCG	۵ ک	н	н
G832u1	WIAF-12577	113387	1129	PAFAHIBI, platelet-activating factor acetylhydrolase, isoform Ib, alpha subunit (45kD)	AGACATTCAC [A/T] GGACACAGAG	E A	E	E-i
G835u1	WIAF-12555	U38276	1311	SEMA3F, sema domain, immunoglobulin domain (Ig), short basic domain, secreted, 3F	CCTCTGGCTC [C/A] GTGTTCCGAG	ر ا	S	တ
G835u2	WIAF-12556	U38276	1229	SEMA3F, sema domain, immunoglobulin domain (Ig), short basic domain, secreted, 3F	ACTCACTITG [A/T] TGAGCTCCAG	T A T	Д	>
G835u3	WIAF-12557	U38276	1473	SEMA3F, sema domain, immunoglobulin domain (Ig), short basic domain, secreted, 3F	GAACCTTCAC [G/A] CCATCTATGA	<u>ل</u> ق	E	E
G835a4	WIAF-13138	U38276	1726	SEMA3F, sema domain, immunoglobulin domain (Ig), short basic domain, secreted, 3F	TGACCAGGAG [A/T] TGGAGGAGCT	E E	Σ	ы
G836u1	WIAF-12592	U28369	1056	SEMA3E, sema domain, immunoglobulin domain (Ig), short basic domain, secreted, 3B	AACGACGTGG [G/A] CGGCCAGCGC	۳ ق ع	ט	Д
G836u2	WIAF-12609	U28369	1479	SEMA3B, sema domain, immunoglobulin domain (Ig), short 1479 basic domain, secreted, 3B	GTCCTGCCCA[C/T]TGGGGGGCGC	<u>ا</u> ت س	F	н

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783811	WTAR-12590	U72671	1107	ICAM5, i	ICAM5, intercellular adhesion	CGCAGCTGGG [A/G] CCCAAGCTCT	Σ	Ą	5	E	ď
				ICAM5, i	intercellular adhesion	חייט גייט סחיים רח' אין מחייט גיטט גיס	۶	٨		<u>-</u>	1
G838u2	WIAF-12591 WTAF-12109	U/26/1 HT961	2232	SOS1, son o	966 Molecule 5, relencepualin SOS1, son of sevenless 2232 (Drosophila) homolog 1	CTCAGGCAAA [T/C] GGAGTAAGCC					· z
1840a2	WTAF-12110	HT961	2404	SOS1, so (Drosophi	SOS1, son of sevenless (Drosophila) homolog 1	ACCGTCTGAA [C/G] TTGTAGGGAG	Σ	υ	U	ı	>
G840u3	WIAF-12213	HT961	3813	SOS1, sc (Drosophi	SOS1, son of sevenless (Drosophila) homolog 1	CAAGGGTACC[G/A]CGTCGATGCT	S	უ	Æ	д	Д
G841u1	WIAF-12153	HT97420	1372	SMOH, sn 1372 homolog	smoothened (Drosophila)	TTTTGGCTTC[C/G]TGGCCTTTGG	М	Ü	ტ	ц	۵
G841u2	WIAF-12179	HT97420	858	SMOH, sn 858 homolog	smoothened (Drosophila)	CCCAGTTCAT [G/T] GATGGTGCCC	Ж	Ð	E-1	Σ	н
G841u3	WIAF-12185	HT97420	1164	SMOH, sn 1164 homolog	smoothened (Drosophila)	CTGTGAGTGG [C/G]ATTTGTTTTG	Ø	ט	Ŋ	_O	ט
G847u1	WIAF-12588	L41939	2019	1	EphB2	GGTCTGCAGT [G/T] GCCACCTGAA	Σ	Ü		ט	U
G847u2	WIAF-12596	L41939	1806	1806 EPHB2, I	EphB2	GTGTAACAGA[A/C]GACGGGGGTT	ω	Ø	ບ		p ₄
G847u3	WIAF-12613	L41939	2885	2885 EPHB2, I	EphB2	AGGCCATCAA [G/C] ATGGGGCAGT	Σ	ro	Ī		z
G848u1	WIAF-12685	L40636	2484	2484 EPHB1, 1	EphB1	GTCAACAGTA[A/G]CCTGGTGTGC	Σ	A		z	co l
G848u2	WIAF-12690	L40636	2020	2020 EPHB1, I	EphB1	CCTTCACTTA [T/C] GAGGATCCCA	တ	E	U	ы	×
G849u1	WIAF-11920	D83492	1544	1544 EPHB6, 1	EphB6	ACCTGTGTGG [C/T] TCATGCAGAG	Σ	ບ	H	Ø	Þ
G849u2	WIAF-11921	D83492	3301	3301 EPHB6,	EphB6	CTTTGGGATA[C/T]TCATGTGGGA	Σ	Ü	н	,그	ſĿ,
G849u3	WIAF-13412	D83492	1139	1139 EPHB6,	EphB6	GAGACCTTCA [C/T] CCTTTACTAC	Σ	ນ	⊢	ы	н
G849u4	WIAF-13413	D83492	1895	1895 EPHB6,	EphB6	TTTGAGGTGC [A/C] AGGCTCAGCA	Σ	A	ບ	oĭ	Д
G849u5	WIAF-13414	D83492	2338	2338 EPHB6,	EphB6	CTATGACCAG [G/A] CAGAAGACGA	Σ	_ت	Æ	Æ	ᇊ
G849u6	WIAF-13415	D83492	2567	2567 EPHB6,	БрћВб	GGGGCTTTGG [C/G] CTTCCTCCTG	Σ	บ		A	Ü
G849u7	WIAF-13422	D83492	2860	2860 EPHB6,	ЕрћВб	GGCCATCCAG [G/A] CCCTGTGGGC	Σ	0	Æ	A	EH
G849u8	WIAF-13423	D83492	2782	2782 EPHB6,	ਤੁਸਪਤ	GGAGGTCATT [G/C] GGACAGGCTC	Σ	r _O	J	U	멊
G849u9	WIAF-13424	D83492	3038	3038 EPHB6,	EphB6	TTCCTCAGGC[A/G]GCGGGAGGGC	Σ	K	ro l	ø	24
G849u10	WIAF-13425	D83492	3637	3637 EPHB6,	БрћВб	AGCCATTGGA[C/T]TGGAGTGCTA	ß	ر ان	H	ᄓ	LI.
G856u1	WIAF-12625	D45906	1323	1323 LIMK2,	LIM domain kinase 2	AGCTGAACCT [G/C] CTGACAGAGT	ß	v	บ	ᄓ	r.
20 1.1.00 1.1.00	MTAT-12630	165019	864		MADH2, MAD (mothers against decapentaplegic, Drosophila) homolog 2	TTTGGTGTTC [G/A] ATAGCATATT	Ŋ	υ	Æ	ഗ	S
G86n1	WIAF-10437	HT1701	263	RAD51, homolog	RAD51 (S. cerevisiae) (E coli RecA homolog)	TGAAGCAAAT [G/C] CAGATACTTC	Œ	_O	Ü	Ą	д
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C11985	WT&R-10465	HT1701	861	RAD51, RAD51 (S. cerevisiae)	GCATCAGCCA [T/C] GATGGTAGAA	Σ	T.	Σ	E+
786113	WIAF-10466	HT1701	924	RAD51, RAD51 (S. cerevisiae) 924 homolog (E coli RecA homolog)	TACAGAACAG [A/G] CTACTCGGGT	Σ			Ŋ
G864a1	WIAF-13139	X82324	183	POU3F4, POU domain, class 3, transcription factor 4	CAGCAATGGG [C/t]ATCCCCTCGG	Σ	U U	н	>-
G866u1	WIAF-12637	HT0101	2576	2576 glutamate receptor (GB:M64752)	AAATCCCGTA [G/A] TGAATCCAAG	×	G B	S	_ z
G866u2	WIAF-12638	HT0101	1131	1131 glutamate receptor (GB:M64752)	TAACAGGAAA [C/T] GTGCAGTTTA	S	H ن	Z	Z
G869u1	WIAF-13406	HT33620	GR. 101 3627 2C	<pre>GRIN2C, glutamate receptor, ionotropic, N-methyl D-aspartate 2C</pre>	AGATCAGCAG [G/T] GTAGCCCGTG	Σ	U U	면	8
G870ul	WIAE-11889	HT4468	714	SLC1Al, solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system 714 Xag), member 1	CAGAAGAGTC [C/G] TTCACAGCTG	S	Ü	හ හ	· · · · · · · · · · · · · · · · · · ·
G870u2	WIAF-11913	HT4468	314	SLCIA1, solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system 314 Xag), member 1	CTAGAGAAT [T/A] CTACTTTGCT	Σ	E-I	F4 (4)	
G870u3	WIAF-11914	HT4468	579	SLCIAl, solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system 579 Xag), member 1	AAGTCAGTAC [G/A] GTGGATGCCA	w o	ပ	E A	E E
G870u4	WIAF-11922	HT4468	706	SLC1Al, solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system 706 Xag), member 1	GAACATGACA [G/A] AAGAGTCCTT	Σ	ъ	A	EI X

G870u5	WIAF-11923	HT4468	876	SLCIA1, solute carrier family 1 (neuronal/epithelial high affinity glutamate transporter, system Xag), member 1	ggaagatcat [a/g] gaagttgaag	<u>ن</u> در ع	н	Σ
G871u1	WIAF-11892	HT3187	1004	SLCIA3, solute carrier family 1 (glial high affinity glutamate transporter), member 3	TTCTCTTAAC [G/C] AAGCCATCAT	ъ Б	H	Ø
G871u2	WIAF-11915	HT3187	1154	SLC1A3, solute carrier family 1 (glial high affinity glutamate transporter), member 3	TGTTGGCTTA [C/T] TCATTCACGC	E E	<u> </u>	ſτι
G871u3	WIAF-11926	HT3187	1412	SLCIA3, solute carrier family 1 (glial high affinity glutamate transporter), member 3	GGCTGCCATT [T/G] TCATTGCTCA	E E	Ĺī4	Δ
G871u4	WIAF-11944	HT3187	1217	SLCIA3, solute carrier family 1 (glial high affinity glutamate 1217 transporter), member 3	AAACCCTIGG [G/A] TTTTATTGG	۳ ع ع	>	Н
1,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	W T D W - 1 3 4 3 3 3	НТ4077	1271	SLCIA2, solute carrier family 1 (glial high affinity glutamate 1271 transporter), member 2	CTGTTGGAGC [A/C] ACCATTAACA	S A C		A
2879111	WTAF-11899	HT28317	1273	GRM2, glutamate receptor,	GACTTTGTGC [T/C] CAACGTCAAG	M H	<u>н</u>	
G879112	WTAF-11932	HT28317	2349	GRM2, glutamate receptor,	CTTCTATGTC [A/G] CCTCCAGTGA	M A G	<u>E</u> +	Æ
G879u3	WIAF-13421	HT28317	2186	GRM2, glutamate receptor, 2186 metabotropic 2	ATGCAAGTAT [G/T] TTGGGCTCGC	E E	Σ	н
G879114	WIAF-13429	HT28317	2567	GRM2, glutamate receptor, 2567 metabotropic 2	CCCAGTITGT [C/T] CCCACTGTTT	S S	٥	>
2879115	WIAF-13436	HT28317	2046	GRM2, glutamate receptor,	ACAGGTGGCC [A/G] TCTGCCTGGC	M A	H	Ν
91182	WTAF-13438	HT28317	2425	GRM2, glutamate receptor, 2425 metabotropic 2	GTGCTTGGCT [G/T] CCTCTTTGCG	Σ Ε	D L	[IL4
G879u7	WIAF-13439	HT28317	2463	GRM2, glutamate receptor, 2463 metabotropic 2	ccrcrrccag[c/T]cgcagaagaa	υ Σ	T.	ω
G880u1	WIAF-12164	HT33719	2117	GRM4, glutamate receptor, 2117 metabotropic 4	AGCCCGACCT[T/G]GGCACCTGCT	ω Ε-Ι	D D	ㅁ

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G880u2	WIAF-12176	HT33719	2427 0	GRM4, glutamate receptor, 2427 metabotropic 4	GGACCTGTCG [C/T] TCATCTGCCT	υ Σ	H	נ	[T4
G880u3	WIAF-12192	HT33719	2372 1	GRM4, glutamate receptor, metabotropic 4	ACCAGCGGAC [A/G] CTCGACCCCC	8	U	E	н
G883a1	WIAF-13140	HT48863	1408	GRM7, glutamate receptor, 1408 metabotropic 7	ATCGCAAATG [C/a] ACAGGACAGG	N N	ď	υ	*
G883a2	WIAF-13141	HT48863	2027	GRM7, glutamate receptor, 2027 metabotropic 7	rccrgrcrrc[c/t]rggcaargrr	ນ ໝ	பு	ᄓ	ı,
G883a3	WIAF-13147	HT48863	1813	GRM7, glutamate receptor, 1813 metabotropic 7	retecacact [a/g] ccatgtaagc	SA	מ	ŀΊ	ᄓ
G883a4	WIAF-13148	HT48863	1536	GRM7, glutamate receptor, 1536 metabotropic 7	TGTGCTGACT[A/t]CCGGGGTGTC	M	t t	>-	Ēτι
G883a5	WIAF-13149	HT48863	2473 1	GRM7, glutamate receptor, 2473 metabotropic 7	AAGCCAGAGG [G/a] GTTCTCAAGT	დ ტ	ಹ	ڻ	r.
G883a6	WIAF-13150	HT48863	2434	GRM7, glutamate receptor, 2434 metabotropic 7	TCATAGACTA[C/t]GATGAACACA	S C	ц	ъ	þ
G884u1	WIAF-11916	U95025	1052	GRM8, glutamate receptor, 1052 metabotropic 8	CGAACTCTTG [C/A] CAATAATCGA	υ Σ	Æ	A	Д
G884u2	WIAF-11945	U95025	2016	GRM8, glutamate receptor, 2016 metabotropic 8	AAACAAACCG [T/C] ATCCACCGAA	S E	υ	<u>R</u>	ద
G884u3	WIAF-11946	095025	1852	GRM8, glutamate receptor, 1852 metabotropic 8	GAGGGCTTCA [G/A] GACGCGAACT	E E	Æ	ರ	ĸ
G884u4	WIAF-11947	095025	2078	GRM8, glutamate receptor, 2078 metabotropic 8	ATTAGTCCAG [C/G] ATCTCAGCTG	Ŭ Σ	ŋ	Æ	U
G884u5	WIAF-13430	U95025	1897	GRM8, glutamate receptor, 1897 metabotropic 8	TTTTCTCTGT [T/G] ATTCAATCAC	E E	U	>-	Д
G884u6	WIAF-13435	U95025	2364	GRM8, glutamate receptor, 2364 metabotropic 8	TTACCATGTA [T/C] ACCACCTGCA	S	υ U	H	×
G885u1	WIAF-13434	AF002700	1363	GFRA2, GDNF family receptor alpha	AACTCAGGCC [C/A] CAGCAGAGCC	Σ	A	Д	н
G886a1	WIAF-13142	U95847	(G) 497 1	GFRA1, GDNF family receptor alpha	GAAGTCGCTC [T/A] ACAACTGCCG	E	ď	>-	Z
G886a2	WIAF-13143	U95847	1385	GFRA1, GDNF family receptor alpha	GTCTGAGAAT [G/a] AAATTCCCAC	Æ	ಹ	四	Ж
G886a3	WIAF-13151	U95847	781	GFRA1, GDNF family receptor alpha	GCGTGTCCAA [T/c] GATGTCTGCA	S	υ	z	z
G892u1	WIAF-11956	U12140	798	NTRK2, neurotrophic tyrosine 798 kinase, receptor, type 2	TGGGCAATCC[A/G]TTTACATGCT	8	<u>.</u>	Д4	
G892u2	WIAF-11957	U12140	834	NTRK2, neurotrophic tyrosine 834 kinase, receptor, type 2	GGATCAAGAC[T/A]CTCCAAGAGG	<u></u>	4	E	E

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U12140
NTRK2, U12140 1738 kinase,
NTRK2, U12140 2486 kinase,
NTRK2, U12140 1106 kinase,
NTRK2, U12140 2085 kinase,
NTRK2, U12140 2230 kinase,
NTRK2, U12140 2223 kinase,
NTRK2, U12140 1602 kinase,
NTRK2, U12140 1354 kinase,
NTRK2, U12140 1944 kinase,
NTRK2, U12140 2103 kinase,
NTRK2, U12140 1860 kinase,
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STTCTGAAAA M T G I M	CACCTAACCT M G T A S	CGCTGGATGC S T C I I	TAGCCCAGCA M A T I L	TTCGTGCACC M C A H Q	GATGCCTCCA N G A W *	GGAACTGCCT S A C R R	TGGTGAAAAT S C T L L	CIGGATAAGG M C T S F	GTTGGTAACC S T C D D	CGCCCTGTCG S C T S S	GGAGGCCTAC M A G Q R	M T C M	S G A	M C A N	SCH	AGTOCOTAN	E . E
GACATAACAT [T/G] GTTCTGAAAA	AAATCTGGCC [G/T] CACCTAACCT	TGCTGCCCAT [T/C] CGCTGGATGC	GATGCTGCAT [A/T] TAGCCCAGCA	CGTCCCAGCA [C/A] TTCGTGCACC	CCCATTCGCT [G/A] GATGCCTCCA	TTTGGCCACC [A/C] GGAACTGCCT	GGAGAACTTG [C/T] TGGTGAAAT	GGGATGTCGT [C/T] CTGGATAAGG	TGTATTGGGA [T/C] GTTGGTAACC	GGTATAAGAG [C/T] CGCCCTGTCG	CCGGAGCTGC [A/G] GGAGGCCTAC	TGCAATTCAA [T/C] GCAGTCCGAA	TGCACACAGT [G/A] GACCACGTGG	ACGTCCGGAA[C/A]AAACTGAAGA	AGCAAGCCCT [C/T] AGTGAGATTG	GCTGAATTAA [G/A] AGTGGCCTAA	C FF FF C C C C C C C C C C C C C C C C
neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	neurotrophic tyrosine receptor, type 2	ferredoxin reductase	ferredoxin reductase	syntaxin 4A (placental)	syntaxin 3A	syntaxin 3A	syntaxin 3A	syntaxin 3A	
NTRK2, 1965 kinase,	NTRK2, kinase,	NTRK2, 2502 kinase,	NTRK2, kinase,	NTRK2, 2364 kinase,	NTRK2, 2507 kinase,	NTRK2, 2389 kinase,	NTRK2, 2416 kinase,	NTRK2, 359 kinase,	NTRK2, kinase,	FDXR,	388 FDXR,	497 STX4A,	758 STX3A,	317 STX3A,	611 STX3A,	909 STX3A,	
N 1965 k	N 958 k	Z502 k	N 2317 k	2364 h	2507 3	2389	2416	359	1044	1130 FDXR,	388	497	758	317	611	606	
U12140	U12140	U12140	U12140	U12140	U12140	U12140	U12140	012140	U12140	J03826	J03826	HT3470	HT27792	HT27792	HT27792	HT27792	
WIAF-13146	WIAF-13442	WIAF-13446	WIAF-13447	WIAF-13448	WIAF-13449	WIAF-13471	WIAF-13472	WIAF-13474	WIAF-13479	WIAF-10222	WIAF-10258	WIAF-11970	WIAF-11969	WIAF-11971	WIAF-12002	WTAF-12003	COORT TOTAL
G892a17	G892u18	G892u19	G892u20	G892u21	G892u22	G892u23	G892u24	G892u25	G892u26	G9u1	G9u2	G900n1	G901u1	G901u2	G901u3	A111000	****

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G901u7	WIAF-13453	HT27792	878	828 STX3A,	3.A		Σ			Į.	Ţ
G901u8	WIAF-13455	HT27792	226	226 STX3A,	3A	IACAGIAICA [1/c] ICICIGCA	E (	Т	T	7	Τ,
G902u1	WIAF-13454	HT27744	848	848 STX5A,	syntaxin 5A	ACTTCCAGTC [T/A] GTCACCTCCA	ς,			T	מ מ
G902u2	WIAF-13456	HT27744	338	338 STX5A,	syntaxin 5A	ATTTCGTGAG [A/G]GCCAAGGGCA	တ	ď	r U	2	×
G905u1	WIAF-12202	HT27789	487	CREBL1,	CAMP responsive element protein-like 1	TCCAGATCAA [C/T] GTTATCCCCA	w	U	H	z	z
G905u2	WIAF-12219	HT27789	151	CREBL1, 151 binding	cAMP responsive element protein-like 1	ATTCTGGCCT [A/T] GATGAAGTGG	တ	Ą	E	- F	Ы
G905u3	WIAF-12230	HT27789	649	CREBL1, 649 binding	cAMP responsive element protein-like 1	AGTCCCTGTC [C/G] CCTTCAGGAT	S	U	U	ω	S
G906u1	WIAF-12214	HT4372	2127	N-ethylı	2127 N-ethylmaleimide-sensitive factor	aagggaagaa [g/a] gtctggatag	ß	ŋ	4	*	M
G906u2	WIAF-12221	HT4372	514	N-ethyl	514 N-ethylmaleimide-sensitive factor	GGGAGAGCCT [G/A] CGACAGGGAA	Σ	U	A.	4	H
G908u1	WIAF-12201	HT3665	98	RAB5A, family	RAB5A, member RAS oncogene	GCCCAAATAC [T/G] GGAAATAAAA	S	ЕН	U	E	E
1,105	WTAF-10438	HT11848	4 0 0	ERCC1, exci complementin deficiency, 1 (includes sequence)	ERCC1, excision repair cross- complementing rodent repair deficiency, complementation group 1 (includes overlapping antisense sequence)	TCGTGCGCAA [C/T] GTGCCCTGGG	ω	Ü	H	Z	Z
Tarke	000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000-000 000 000-000 000 000 000-000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 000 00	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0			ERCC1, excision repair cross- complementing rodent repair deficiency, complementation group 1 (includes overlapping antisense	CTRAGAGCCAC IG /A ] TGCCCCACAG	S	Ů	A.	E	H
G91u2	WIAF-13210	HT3672	252	synapto	synaptobrevin 1	GCAGTGCTGC [C/A] AAGCTAAAGA	S	U	Æ	4	A
1 67 LOG	WTAF-12115	D63506	1390		Homo sapiens mRNA for unc- 18homologue, complete cds.	TTACCTTGGT [G/A] TTCCCATTGT	Σ	Ö	A	۸	н
2915112	WIAF-12293	D63506	685	Homo sapiens	apiens mRNA for unc- logue, complete cds.	ACAGCTTGTT [G/A]AAAAAAGCT	Σ	ტ	Æ	ш	×
1691691	WTAF-13209	HT28523	308	Huntingtin a	Huntingtin associated protein 1- like protein	GAGCAGTITT [C/T] GGAGGCCAGC	Σ	บ	EH	Ø	д
2631692	WIAF-13211	HT28523	762	Hunti like	ngtin associated protein 1- protein	CGGAGGAGTT [G/C] GTGCCCCAGG	Σ	<u></u> છ	บ	ц	ſъ
G916a3	WIAF-13212	HT28523	56(	Huntingtin a 560 like protein	Huntingtin associated protein 1- like protein	GAGCTCAGAA [C/T] GTCTCTAAGG	Σ	<u></u> υ	H	F	×

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	WTAF-11972	1179734	1075 0	HIP1, hu 1075 protein 1	ntingtin interacting	AGAGCCAGCG [G/A] GTTGTGCTGC	S S	ტ	Æ	요 요	_
2917112	WTAF-11973	U79734	1005 p	HIP1, hu protein 1	ntingtin interacting	GACCACTTAA [T/C] TGAGCGACTA	A'	E-4	ບ	H	
	WIAF-11977	U79734	1539 p	HIP1, hu 539 protein 1	ntingtin interacting	CTGCAAGGCA [G/A] CCTGGAAACT	H.	Ŋ	4.	S	
	WTAF-12005	1179734	817 g	HIP1, h	ntingtin interacting	TGGTGGTGAT [C/T] CCTGCAGAGG	S	ນ	EH	I	
	WTAF-12006	1179734	19061	HIP1, h	huntingtin interacting	GCTGGAGCCA [G/C] TATCTGGCCT	T. M	D.	U	н ŏ	
	WIAF-13157	U79734	993 E	HIP1, hu protein 1	ntingtin interacting	aaggatgaga [a/g]ggaccactta	TA M	Ą	ט	저	
G919u1	WIAE-11974	D30742	707	CAMK4, dependen	CAMK4, calcium/calmodulin-	ACTGCGCACC [T/C] GAAATTCTTA	মূ	EH	υ	<u>д</u>	
G919u2	WIAF-11991	D30742	1139	CAMK4, dependen	CAMK4, calcium/calmodulin- 1139 dependent protein kinase IV	AGAGCCACAA [G/A] GCTAGCCGAG	S S		Æ	X X	
G919u3	WIAF-12007	D30742	834	CAMK4, dependen	CAMK4, calcium/calmodulin-834 dependent protein kinase IV	CATGTTCAGG [A/T] GAATTCTGAA	N	A.	E-I	*	
710100	MTAE-12443	n30742	1088	CAMK4, depender	CAMK4, calcium/calmodulin-	TGGCCTCTTC[C/G]CGCCTGGGAA	s s	ပ	G	S S	
G92011	WIAF-11979	X78520	1952	1952 CLCN3,		ATGACATTCC [T/C] GATCGTCCAG	A.G. S.	E	Ü		
G920u2	WIAF-11980	X78520	1819	1819 CLCN3,	chloride channel 3	ATAGCCTTCC [C/T] TAATCCATAC			н		Ţ
G920u3	WIAF-11981	X78520	2094	2094 CLCN3,	chloride channel 3	CATTGGAGCG [A/G] TCGCAGGAAG			r.		
G920u4	WIAF-11983	X78520	2822	2822 CLCN3,	chloride channel 3	ATATTTTCCG [A/G] AAGCTGGGAC			_D		p4
G920u5	WIAF-11984	X78520	2745	2745 CLCN3,	channel 3	GCCATTGAAG [C/T] TTCGAAGCAT			E→ {		[zı [
G920u6	WIAF-11987	X78520	2499	2499 CLCN3,	3	TCCCTTAGCT [G/T] TCCTGACACA			<u> </u>		±, 0
G920u7	WIAF-12008	X78520	1251	1251 CLCN3,	channel 3	CATCATCAGA [G/A] GTTACTIGGG			₹ E		ן מ
G920u8	WIAF-12011	X78520	888	888 CLCN3,	channel 3	AGTAGTAACA [C/T] TAACAGGATT			:   ნ	] ]	<b>⊐</b> ⊦
G920u9	WIAF-13459	X78520	2804	2804 CLCN3,	chloride channel 3	CAATGGAGAT [T/C] GTGGTGGATA	A L	-	ر		
				CLU, clust inhibitor, glycoprote:	CLU, clusterin (complement lysis inhibitor, SP-40,40, sulfated glycoprotein 2, testosterone-						
G921u1	WIAF-11954	J02908	931	repress apolipo	repressed prostate message 2, 931 apolipoprotein J)	GAGAGGTTGA [C/T] CAGGAAATAC	'AC M	<u>ပ</u>	H	[+	н

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CCCTCCCAGG [C/T] TAAGCTGCGG	CTCACGCAAG [G/C] CGAAGACCAG	TCAACACCTC [C/T] TCCTTGCTGG	GAGCTAAGCC [G/A] GGGCAAGCTC	CTAAGCCGGG [G/T] CAAGCTCTAT		GGGTCATGAG [T/C] GTCTGTCTGC	TGCTGCCCAT [C/T] CGCTGGATGG	TACCAGGAGC [C/T] CCGGCCTCGT	CGCCCCACTC [C/T] GCTCCCTGTG	TGGAGAACGG [C/T] GACCTCAACC	TGAAAGCTTT [G/T] ACCTGGAGCC	CCACGCGATT[C/G]ATCAGGATCT	GACCTTCTGG [T/C] ATCACATGTC
CLU, clusterin (complement lysis inhibitor, SP-40,40, sulfated glycoprotein 2, testosteronerepressed prostate message 2, 880 apolipoprotein J)	CLU, clusterin (complement lysis inhibitor, SP-40,40, sulfated glycoprotein 2, testosteronerepressed prostate message 2, 1051 apolipoprotein J)	CLU, clusterin (complement lysis inhibitor, SP-40,40, sulfated glycoprotein 2, testosteronerepressed prostate message 2, 986 apolipoprotein J)	Human 2',3'-cyclic nucleotide 3'- phosphodiesterase mRNA, complete cds.	Human 2',3'-cyclic nucleotide 3'- phosphodiesterase mRNA, complete 1062 cds.	Human 2',3'-cyclic nucleotide 3'- phosphodiesterase mRNA, complete 1141 cds.	cell	cell adhesion	1601 Cak, cell addesion Almase	cell	cell	577 NRP1, neuropilin 1	1683 NRP1, neuropilin 1	2176 NRP1, neuropilin 1
102908	J02908	J02908	M19650	0.0				1,11315			AF018956	AF018956	AF018956
WIAF-11955	WIAF-11990	WIAF-13469	WIAF-11993	WTAR-11994	WIAR-13445			WIAF-11996	WIRE-13440	WIAF-13451	WIAF-11961	WIAF-11963	WIAF-11975
G921u2	G921u3	G921u4	G923u1	64503n0	G923u3	G925u1	G925u2	G925u3	G92544	925116	G926ul	G926u2	G926u3

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G926u4         WIAF-           G926a5         WIAF-           G926a6         WIAF-           G926u7         WIAF-           G926u8         WIAF-           G926u9         WIAF-           G926u10         WIAF-           G927u1         WIAF-	WIAF-11976 WIAF-13158	AF018956	2092 NRP1		П	TTCCCAAGCT [G/T] AUGAAATUA	Ξ		1 H	ı E	T
0	-13158	7000 1011	747 NRP1			100000000000000000000000000000000000000	C				
0		AFULGYDD				TTTTTTACAC [C/T] GACAGCGCGA	ν (	T	Τ	Ĺ	Т
0	WIAF-13159	AF018956	996 NRP1		F-1	ACTTGGGCCT [T/C] CTGCGCTTTG	מ :				Т
0	WIAF-13444	AF018956	644 NRP1		1	GAAATCTGGG [A/C] TGGATTCCCT	Z				
	WIAF-13450	AF018956	1738 NRP1		neuropilin 1	CAGAATGGAG [C/G] TGCTGGGCTG	Σ				_
	WIAF-13452	AF018956	537 NRP1		neuropilin 1	TIGICITIGC [G/A] CCAAAGAIGI	တ				Т
	WIAF-13457	AF018956	2197 NRP1		neuropilin 1	TGGGTCCCAC [G/A] TCGGCACACT	Σ				
	WIAF-11978	AF022860	870 NRP2		neuropilin 2	GGATTGCTAA [T/C] GAACAGATCA	ß	T			Т
	WIAF-11982	AF022860	1674 NRP2,		neuropilin 2	ATGACACCCC [T/G] GACATCCGAA	S				Т
	WIAF-11985	AF022860	1250 NRP2,		neuropilin 2	TGGCACTCAG [G/A] TATCGCCCTC	Σ				$\neg$
	WIAF-11986	AF022860	1071 NRP2		neuropilin 2	ATGGCTACTA[C/T]GTCAAATCCT	ഗ				
	WIAF-12009	AF022860	726 NRP2,	١.	neuropilin 2	GTTCATCGAC [G/A] GGGATCCTCT	ഗ			$\neg$	T
G927u6 WIAF	WIAF-12010	AF022860	2522 NRP2		neuropilin 2	GCAACCTCAG [G/T] GTCTGGCGCC	Σ				T
	WIAF-12012	AF022860	123 NRP2		neuropilin 2	GCTATATCAC [C/T] TCTCCCGGTT	S	$\neg$			
	WIAF-13160	AF022860	2427 NRP2,		neuropilin 2	CTTTTGCAGT [G/T] GACATCCCAG	ß				
	WIAF-13161	AF022860	2430 NRP2		neuropilin 2	TTGCAGTGGA [C/G] ATCCCAGAAA	Σ	บ	r.	D E	
	WIAF-13162	AF022860	2463 NRP2,		neuropilin 2	AAGGATATGA [A/G]GATGAAATTG	ß	Æ			
	WIAF-13163	AF022860	2473 NRP2		neuropilin 2	AGATGAAATT [G/T] ATGATGAATA	Σ	Ö	H	<del>Х</del> Д	
	WIAF-13480	AF022860	724 NRP2		neuropilin 2	TCGTTCATCG [A/T] CGGGGATCCT	Σ	Æ	- 1	T.	
	WIAF-13481	AF022860	767 NRP2		neuropilin 2	ATGGCGGTGG [C/T] CAAGGATGGC	Σ	υ	E	A	
	WIAF-13164	HT2608	GABRA2 609 (GABA)	্ৰ	gamma-aminobutyric acid receptor, alpha 2	acaatgggaa [g/a] aaatcagtag	တ	ro l	ಹ	×	×
	WIAF-13153	HT2609	GABRA3 1111 (GABA)	` ₫	gamma-aminobutyric acid receptor, alpha 3	actggttcat [a/g] gccgtctgtt	Σ	ď	מ	Н	Σ
	WIAF-13165	HT2609	GABRA3	্ৰ	gamma-aminobutyric acid receptor, alpha 3	TGTCAGCAAG[G/A] TTGACAAAAT	Σ	Ŋ	Ą	>	н
	WIAF-13154	HT27773	GAB 1077 (GA	GABRA4, ge (GABA) A re	gamma-aminobutyric acid receptor, alpha 4	caaaagaag [a/g] catcaaagcc	Σ	Æ	_O	L L	A
	WIAF-13155	HT27773	GAB 1189 (GA	GABRA4, ge (GABA) A re	gamma-aminobutyric acid receptor, alpha 4	AGAACAAATG [C/A] TTTGGTTCAC	Σ	Ü	Æ	A	А
G936ul WIAE	WIAF-12308	HT3432	GAB 1027 (GA	GABRB2, ga (GABA) A re	gamma-aminobutyric acid receptor, beta 2	AATTACGATG [C/T] TTCAGCTGCA	Σ	υ	E+	Æ	<b>A</b>
	WIAF-12327	HT3432	GABRB2   362 (GABA)	` ≰	gamma-aminobutyric acid receptor, beta 2	AAGGCTATGA [C/T] ATTCGTCTGA	Ω	υ	E	Д	Д



IGATACCTAT M C T P L	CTACAGTGAG M G C S T	CACGGGCGTG M T C I T	TCAAGGCCGT M G A V I	GITCACCACT M A C Q P	GCCACATTTG M A C K T	AACGIGCAAG M C I A V	ACCGICCAGA S T C F F	CAGAAACTAC M C A P T	AGTCCGAGCC M G a B K	CATGCTCTGT M A 9 Y C	GAGTITAAAA S T C D D	GCCAACATCA S C t N N	0.00
CTCTGGGTGC[C/T]TGATACCTAT	CTGGATGGAA [G/C] CTACAGTGAG	ACCACCATCA [T/C] CACGGGCGTG	1 CGTCTCCTAC [G/A] TCAAGGCCGT	GTCCTGCTCC [A/C] GTTCACCACT	GATAACAGCA [A/C] GCCACATTG	CTGGGTAGTG[C/T]AACGTGCAAG	CTGGCCTCTT [T/c] ACCGTGGAGA	CTACCCCAAC [C/a] CAGAAACTAC	GTGTGCCCCA [G/a] AGTCCGAGCC	ATCAGCTTCT [A/g] CATGCTCTGT	ACCACCTGGA [T/c] GAGITIAAAA	CCGGCTCCAA[C/t]GCCAACATCA	
GABRB2, gamma-aminobutyric acid 571 (GABA) A receptor, beta 2	GABRR2, gamma-aminobutyric acid (GABA) receptor, rho 2	GABRR2, gamma-aminobutyric acid (GABA) receptor, rho 2	GABRR2, gamma-aminobutyric acid (GABA) receptor, rho 2	Human putative G protein-coupled receptor (GPR19) gene, complete ods.	Human putative G protein-coupled receptor (GPR19) gene, complete cds.	Human putative G protein-coupled receptor (GPR19) gene, complete cds.	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. transcript 1	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. transcript 1	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. transcript 1	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. 5703 transcript 1	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. 5809 transcript 1	calcium channel, voltage-gated, alpha 1 subunit, L type, alt. 6616 transcript 1	
2 571	6 1219	6 1003	G HT2236 1041	Huma rece U64871 785 cds.	443	71 818	5110	3842	5624	5703	5809	60 6616	
WIAF-12328 HT343	WIAF-12330 HT223	WIAF-12355 HT223	WIAF-12356 HT2			WIAF-13625 U648							
G936u3	G939u1	G939u2	G939u3	205011	200000	95013	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	1 C	м м м м м м м м м м м м м м м м м м м	00000000000000000000000000000000000000	יין מין מין מין מין מין מין מין מין מין	995586	

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G956u2	WIAF-14188	HT2199	1452	calcium channel, voltage-gated, 1452 alpha 1D subunit, DHP-sensitive	AAGAGGACCC [A/T]GCTCCATGTG	8 8	H	Д	д
G956u3	WIAF-14189	HT2199	1614	calcium channel, voltage-gated, 1614 alpha 1D subunit, DHP-sensitive	GCTGGACAGA[C/T]GTGCTCTACT	ა ე	H	Д	D
G956u4	WIAF-14190	HT2199	2540	calcium channel, voltage-gated, 2540 alpha 1D subunit, DHP-sensitive	GGCAAGITIA [A/I] TITIGAIGAA	M A	E	z	н
G956u5	WIAF-14191	HT2199	3210	calcium channel, voltage-gated, 3210 alpha 1D subunit, DHP-sensitive	TGCTGAGCAG [T/C] GCTGCCCTGG	S	٢	တ	S
9n926Ð	WIAF-14192	HT2199	3326	calcium channel, voltage-gated, 3326 alpha 1D subunit, DHP-sensitive	TTGAAGATGA[C/T]AACTTTTGGA	υ Σ	E	E-	н
G956u7	WIAF-14193	HT2199	3274	calcium channel, voltage-gated, alpha 1D subunit, DHP-sensitive	ACTGGGTTAC [T/C] TTGACTATGC	E	U	[II4	
6956u8	WIAF-14194	HT2199	5127	calcium channel, voltage-gated, 5127 alpha 1D subunit, DHP-sensitive	TGCCTCTCAA [C/T] AGTGACGGGA	S C	E-4	Z	z
6956b	WIAF-14195	HT2199	5173	calcium channel, voltage-gated, alpha 1D subunit, DHP-sensitive	TGCTTTGGTT [C/T] GAACGGCTCT	Z U	E	<u>µ</u>	*
G956u10	WIAF-14200	HT2199	1437	calcium channel, voltage-gated, 1437 alpha 1D subunit, DHP-sensitive	CAGATATCGT [A/G] GCTGAAGAGG	\cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot	<u> </u>	>	>
G956ull	WIAF-14201	HT2199	2567	calcium channel, voltage-gated, 2567 alpha 1D subunit, DHP-sensitive	ACCAAGCGGA [G/T] CACCTTTGAC	Æ	E-1	<u> </u>	н
G956u12	WIAF-14202	HT2199	4464	calcium channel, voltage-gated, 464 alpha 1D subunit, DHP-sensitive	TCACCTTTTT [C/T] CGTCTTTTCC	8	E	(Sty	[tr
G956u13	WIAF-14215	HT2199	6927	calcium channel, voltage-gated, alpha 1D subunit, DHP-sensitive	GCTACAGCGA [C/T] GAAGAGCCAG	8	E-1	Ð	
G956u14	WIAF-14216	HT2199	6858	calcium channel, voltage-gated, 6858 alpha 1D subunit, DHP-sensitive	CCCGAGCCAA[C/T]GGGGATGTGG	S C	E-I	Z	Z
G957u1	WIAF-12306	HT4229	915	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	TACATCGAGC [G/A] TGCTTCATGA	<u>ن</u> ک	A	· ·	<u>¤</u>

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G957u2	WIAF-12309	HT4229	3555	alpha 1E subunit, alt. transcript 2	GCCACTACAT [C/T] GTGAACCTGC	S C	H	н	н
G957u3	WIAF-12310	HT4229	4116	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	ATGTAGATCA [C/T] GAGAAAACA	ა ა	E	д	H
G957u4	WIAF-12313	HT4229	5181	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	agaacgagaa [T/C] gaacgctgcg	S	ت ت	z	Z
G957u5	WIAF-12314	HT4229	5971	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	TATGGACCCC [G/A] CCGATGACGG	<u>.</u> დ	Ą	E	H
G957u6	WIAF-12315	HT4229	5985	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	ATGACGGACA [G/T] TTCCAAGAAC	<u>ت</u>	Ð	Ø	н
G957u7	WIAF-12329	HT4229	3100	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	GCTGGCAGGA [G/A] GCCTTGATGA	<u>ნ</u>	4	r D	S
G957u8	WIAF-12331	HT4229	6492 2 a. 2	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	CCCTCCTTTC [C/T] TACAGCTCCC	<u>Σ</u>	F	٥٠	ద
G957u9	WIAF-12354	HT4229	3839 239 20 20 20 20 20 20 20 20 20 20 20 20 20	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	aacgctttgg [g/c] aaccaacaa	<u>ත</u>	ပ	ტ	ď
G957u10	WIAF-12357	HT4229	4753	calcium channel, voltage-gated, alpha 1E subunit, alt. transcript 2	TGACTTCATC [A/G] CCGTGATTGG	M	უ	E	Ą
G960u1	WIAF-12305	HT3336	1246	CACNB3, calcium channel, voltage- 1246 dependent, beta 3 subunit	TTGATGCCCT [C/T] TGATGAGGCC	υ Σ	H	ω	[14
G960u2	WIAF-12340	HT3336	1288	CACNB3, calcium channel, voltage- 1288 dependent, beta 3 subunit	TGGACAGGAT [C/T] TTCACAGCGT	Σ Σ	<del>[+</del>		<u> </u>
6960u3	WIAF-12345	HT3336	641	CACNB3, calcium channel, voltage- 641 dependent, beta 3 subunit	AGGCTCTCTT [C/T] GACTTCCTCA	S C	E		[St.)
G960u4	WIAF-12346	HT3336	576	CACNB3, calcium channel, voltage- 576 dependent, beta 3 subunit	CATGCGGCCT [G/A] TGGTGCTGGT	Σ Σ	4.	>	Σ
G961u1	WIAF-12322	U95019	2037	CACNE2, calcium channel, voltage-	ACTCTGCCTA[C/T]GTAGAGCCAA	<u>s</u>	H	*	×

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G961u2	WIAF-12347	U95019	2007	CACNB2, calcium channel, voltage-	CATTIGACTC [G/A] GAAACCCAGG	r U	8	Ω
G962u1	WIAF-12324	U95020	1423	CACNB4, calcium channel, voltage-	CCAATTGAAA[G/A]ACGAAGTCTA	5	A R	×
G962u2	WIAF-12342	U95020	167	CACNB4, calcium channel, voltage- 167 dependent, beta 4 subunit	GGAGCAGGTT [G/T] AAAAGATCCG	უ	H H	[74
G962u3	WIAF-12350	095020	1571	CACNB4, calcium channel, voltage- dependent, beta 4 subunit	ACACTTACAA [A/G] CCCCATAGGA S	A.	ρ ×	<u>×</u>
G965u1	WIAF-12312	040583	1276	CHRNA7, cholinergic receptor,	TCCTGCACGG [T/C] GGGCAACCCC	H	ڻ ن	r to
G968a1	WIAF-12119	HT27592	1008	CHRNAl, cholinergic receptor, nicotinic, alpha polypeptide 1 (muscle)	ACACACACCA [C/T] CGCTCACCCA	υ	H	田
G968u2	WIAF-12368	HT27592	1136	CHRNA1, cholinergic receptor, nicotinic, alpha polypeptide 1 (muscle)	AAGATTTTTA[C/T]AGAAGACATT M	υ	E E	н
G973a1	WIAF-13172	HT48774	800	CHRNA2, cholinergic receptor, nicotinic, alpha polypeptide 2 (neuronal)	acacticaga [c/t] giggigatig s	ບ	Ω L	<u>P</u>
G973a2	WIAF-13173	HT48774	927	CHRNA2, cholinergic receptor, nicotinic, alpha polypeptide 2 (neuronal)	CTGGAACCCC[G/a] CTGATTTTGG M	უ	ر لا	Ħ
G977u1	WIAF-13949	Y08419	366	CHRNAS, cholinergic receptor, 366 nicotinic, alpha polypeptide 5	AAGTTATACG [T/C] GTTCCTTCAG	E	C R	~ ~
G978a1	WIAF-13179	Y08417	1331	CHRNB3, cholinergic receptor, nicotinic, beta polypeptide 3	CCATTAGATA [C/a] ATTTCGAGAC N	υ	B	
G983a1	WIAF-13214	HT0374	236	236 NPY, neuropeptide Y				-
G983a2	WIAF-13215	HT0374	290	290 NPY, neuropeptide Y				
G983a3	WIAF-13216	HT0374	111	111 NPY, neuropeptide Y	GCGACTGGGG [C/T] TGTCCGGACT S	ט	E I	긔
G987al	WIAF-13174	HT27830	159	PPYR1, pancreatic polypeptide 159 receptor 1	TGGTCTTCAT [C/T] GTCACTTCCT S	υ	ь	н
G987a2	WIAF-13175	HT27830	222	PPYR1, pancreatic polypeptide receptor 1	TGATGTGTGT [G/A] ACTGTGAGGC	Ŋ	Æ	<u>N</u>
G987a3	WIAF-13176	HT27830	322	PPYR1, pancreatic polypeptide 322 receptor 1	GCCGCTGACC [G/T] CCGTCTACAC	ტ	E	8

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G987a4	WIAF-13177	HT27830	1074	PPYR1, pancreatic polypeptide.	TGGAGGAGTC [G/A] GAGCATCTGC	۵ ا	တ	ω
G987a5	WIAF-13178	HT27830	975	PPYR1, pancreatic polypeptide 975 receptor 1	CCTCCACCTG [C/T] GTCAACCCAT S	D D	บ	Ü
G987a6	WIAF-13180	HT27830	615	PPYR1, pancreatic polypeptide 615 receptor 1	AGTTCCTGGC [A/g] GATAAGGTGG	Ą	4	K
G987a7	WIAF-13181	HT27830	718	PPYR1, pancreatic polypeptide 718 receptor 1	GGGCTTCATC [C/T] TGGTCTGTTA S	H ت	Ľ	ъ
G987a8	WIAF-13182	HT27830	745	PPYR1, pancreatic polypeptide 745 receptor 1	CATCTACCGG [C/t] GCCTGCAGAG	Ω t	ద	U
G987a9	WIAF-13183	HT27830	842	PPYR1, pancreatic polypeptide 842 receptor 1	GTGATGGTGG [T/A] GGCCTTTGCC M	F F	>	田
G987a10	WIAF-13184	HT27830	852	PPYR1, pancreatic polypeptide receptor 1	TGGCCTTTGC[C/T]GTGCTCTGGC	ط ن	4	A.
G987a11	WIAF-13185	HT27830	88 89	PPYR1, pancreatic polypeptide receptor 1	CAACAGCCIG [G/a] AAGACIGGCA M	т С	田	×
G987a12	WIAF-13186	HT27830	924	PPYR1, pancreatic polypeptide receptor 1	CCATCTGCCA [C/T] GGGAACCTCA	D D	Ħ	н
G989u1	WIAF-13573	D86519	891	891 NPY6R, neuropeptide Y receptor Y6	Y6 TGACTCATGC[C/T]TACTGGGGCA	H ن	Æ	A
G989u2	WIAF-13588	D86519	465	465 NPY6R, neuropeptide Y receptor Y6	receptor Y6 ACCACCCAGC[A/G]TCTAATACAA S	A G	Æ	Ą
G989u3	WIAF-13591	D86519	980	980 NPY6R, neuropeptide Y receptor Y6	Y6 GAGCCCTTCC [G/A] CAACCTCTCT M	ව අ	_α	Ħ
G991u1	WIAF-12390	HT97376	336	336 Notch2	AAGGTACTTG [C/T] GTTCAGAAAA S	C	υ	ပ
7,000	WTAF-12359	1195299	1343	NOTCH4, Notch (Drosophila)	TCCACACTCT [G/T] CCTGTGTCAG	D D	υ	[îz _i
C112005	WTAF-12361	U95299	2020	NOTCH4, Notch (Drosophila)	TAAGGACCAG [A/G] AAGACAAGGC	A G	×	Ĺτj
G993u3	WIAF-12384	U95299	5775	NOTCH4, Notch (Drosophila) 5775 homolog 4	GGGCCTATTC [G/T] CATTGCCGGA	D	ß	Ø
G996a1	WIAF-13213	HT3329	356	356 OPRM1, opioid receptor, mu 1	CTTAGATGGC [A/G] ACCTGTCCGA	Ą	z	Д
LPLa4	WIAF-13314	HT1320	443	443 LPL, lipoprotein lipase				н
LPLa5	WIAF-13315	HT1320	579	579 LPL, lipoprotein lipase				D.
LPLa6	WIAF-13316	HT1320	609	609 LPL, lipoprotein lipase				ы
LPLa7	WIAF-13317	HT1320	1338	LPL, lipoprotein lipase				E
LPLa8	WIAF-13318	HT1320	1117	lipoprotein		E C	>	<b>Ω</b> [
гРга9	WIAF-13319	HT1320	715	lipoprotein				ם מ
LPLa10	WIAF-13320	HT1320	834	lipoprotein	CTGGTCGAG [C/A] ATTGGAGTCC M	ע _ב	מ כ	χ [ <del>χ</del>
LPLa11	WIAF-13321	HT1320	100 L			ر .		1 *
LPLa12	WIAF-13322	HT1320	1595	595 LPL, lipoprotein lipase		,		]

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LPLa13	WIAF-13323	HT1320	1597 LPL,	lipoprotein lipase	TAAGAAGTCA [G/A] GCTGAAACTG	Σ Ω	Ø	Ŋ
LPLa14	WIAF-13324	HT1320	1606 LPL,	lipoprotein lipase	AGGCTGAAAC [T/C] GGGCGAATCT	_ T	บ	
LPLa15	WIAF-13325	HT1320	1611 LPL,	lipoprotein lipase	GAAACTGGGC [G/A] AATCTACAGA	- G	A	

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While this invention has been particularly shown and described with references to preferred embodiments thereof, it will be understood by those skilled in the art that various changes in form and details may be made therein without departing from the scope of the invention encompassed by the appended claims.